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- (71) Applicant (for all designated States except US): EURO-CELTIQUE S.A. [LU/LU]; 122, Boulevard de la Petrusse, L-2330 Luxembourg (LU).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): KAMMESHEIDT, Anja [DE/US]; 31558 Eagle Rock Way, Laguna Beach, CA 92651 (US). HODGES, Dianne [US/US]; 14351 Pinewood Road, Tustin, CA 92780 (US).
- (74) Agents: ROBINSON, Joseph, R. et al.; Darby & Darby P.C., P.O. Box 5257, New York, NY 10150-5257 (US).

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(54) Title: SPLICE VARIANT OF HUMAN SODIUM III CHANNEL (HNAII118)

(57) Abstract: Described herein is a splice variant of the human NaIII channel α subunit, designated hNaIII18. Also described are nucleotide and amino acid sequence for hNaIII18, oligonucleotide primers and probes for hNaIII18, hNaIII18 regulatory sequences, hNaIII18-specific antibodies, methods of detecting hNaIII18 proteins or nucleic acids, and methods of screening for modulators of hNaIII18 expression or activity.

# Splice Variant of Human Sodium III Channel (hNaIII18)

This application claims priority from U.S. Provisional Application Serial No. 60/431,794, filed December 4, 2002, which is hereby incorporated by reference in its entirety.

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#### FIELD OF THE INVENTION

The present invention relates to a human splice variant of the voltage-gated sodium III channel, termed hNaIII18, as well as methods for stable expression of hNaIII18 in cell lines, and methods of use in screening for compounds that modulate sodium channel activity.

## BACKGROUND OF THE INVENTION

Sodium channels are voltage-gated transmembrane proteins that are involved in the generation of action potentials in electrically excitable cells such as neurons and muscle cells. They are responsible for the cellular uptake of sodium during electrical signals in cell membranes. The channels are members of a multigene family of transmembrane proteins and are typically composed of a large transmembrane pore-forming  $\alpha$ -subunit and three smaller accessory  $\beta$ -subunits (Cattrall et al., Adv Neurol 1999; 79:441-56). The primary structure of  $\alpha$ -subunits is conserved among different sub-types and species. The  $\alpha$ -subunit is all that is required for the channel to be fully functional, however, the  $\beta$ -subunits have been shown to modulate the function of the channel. Specifically, co-expression of rat  $\beta$ 1,  $\beta$ 2, and  $\beta$ 3 subunits with the Na(v)1.2a  $\alpha$ -subunits in the tsA-201 sub-clone of HEK293 cells shifted sodium channel activation and inactivation to more positive membrane potentials. The  $\beta$ 3 subunit alone caused increased persistent sodium currents. (Qu et al., Mol Cell Neurosci 2001;18(5):570-80).

Previous studies have demonstrated numerous different types of  $\alpha$ subunits, which are categorized based on their sensitivity to tetrodotoxin (a toxin
produced by the puffer or fugu fish). Subunits that are inhibited by nanomolar
concentrations of tetrodotoxin are generally referred to as tetrodotoxin-sensitive
channels (TTX-S), while those that require at least micromolar concentrations for
inhibition are referred to as tetrodotoxin-resistant channels (TTX-R).

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Rapid entry of sodium ions into cells causes depolarization and generation of the action potential. Such entry of sodium ions through sodium channels in response to a voltage change on the plasma membrane in excitable cells plays a functional role in control of neuronal excitability in the central nervous system (CNS) and peripheral nervous system (PNS).

An increase in the rate of spontaneous firing in neurons is often observed in peripheral sensory ganglia following nerve injury (Ochoa and Torebjork, Brain1980; 103(4):835-53.; Nordin et al., Pain 1984; 20(3):231-45; Matzner et al., J Neurophysiol 1994;72(1):349-59; Woolf, Drugs 1994; 47 Suppl 5:1-9; discussion 46-7). It has been suggested that this hyperexcitability in neurons is due to altered sodium channel expression in some chronic pain syndromes (Tanaka et al., Neuroreport 1998; 9(6):967-72). Increased numbers of sodium channels leading to inappropriate, repetitive firing of the neurons have been reported in the tips of injured axons in various peripheral nervous tissues such as the DRG, which relay signals from the peripheral receptors to the central nervous system (Waxman and Brill, Biophys J 1978; 21(2):147-60; Devor et al., Neurosci Lett 1989;102(2-3):149-54; Matzner and Devor, Brain Res 1992; 597(1):92-98). Transcripts encoding the  $\alpha$ III subunit, which are present at only very low levels in control DRG neurons, are expressed at moderate to high levels in axotomized DRG neurons together with elevated levels of  $\alpha I$  and  $\alpha II$ mRNAs (Waxman et al, Brain Res Mol Brain Res 1994; 22(1-4):275-89). Conversely, transcripts of sodium channel  $\alpha$  subnits in the sensory nervous system are down-regulated in DRG neurons following axotomy (Dib-Hajj et al., Proc Natl Acad Sci U S A. 1996; 93(25):14950-4). Furthermore, the partial efficacy of sodium blocking agents is well documented in patients treated for neuropathic pain (Omana-Zapata et al., Pain 1997; 7 2(1-2):41-9; Rizzo, J Neurophysiol 1997; 77(1):236-46), providing an important link between increased sodium channel expression and

neuropathic pain. Therefore, alterations in sodium channel expression and subsequent function may be a key molecular event underlying the pathophysiology of pain after peripheral nerve injury.

The partial type III isoform (α-subunit) of the human sodium channel gene, SCN3A, isolated from placenta, was first described by Malo et al. (Proc Natl Acad Sci U SA 1994; 91(8):2975-9; GenBank Accession No. S69887). Two alternative isoforms, neonatal and adult forms, of SCN3A were thereafter identified in human brain tissue by Lu and Brown (J Mol Neurosci 1998;10(1):67-70; GenBank Accession Nos. AF035685 and AF035686, respectively). These isoforms contained a 92 amino acid insert within a region containing putative splice sites (identified through sequence homology with the rat type III brain sequence). The complete coding sequences for human SCN3A genomic DNA and mRNA (and the corresponding protein sequence) also cloned from human brain, was described by Clare et al. (Ann NY Acad Sci. 1999;868:80-3; GenBank Accession Nos. AJ251507 (SEQ ID NO: 3-Figure 3) and AF225987 (SEQ ID NO: 4-Figure 4, respectively).

Most recently, in 2000, Jeong et al. submitted to GenBank an mRNA sequence encoding a splice variant of human SCN3A (Accession No. AF225987; SEQ ID NO: 5-Figure 5). The amino acid sequence of this splice variant contained a 49-amino acid insert from residues 624 to 673 (SEQ ID NO: 6 - Figure 6), when compared with the sequence described by Clare et al. (*supra*).

There remains a need in the art to identify and characterize additional human sodium channels and variants thereof, in order to assist in the identification of drug candidates that can be used to treat conditions involving or associated with over-or under-expression, or over- or under-activated sodium channels.

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#### SUMMARY OF THE INVENTION

The present invention provides a novel splice variant of human sodium channel III  $\alpha$  subunit, designated herein as "hNaIII18", having the amino acid sequence of SEQ ID NO: 2 (Figure 2).

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The present application also provides an isolated nucleic acid having a nucleotide sequence encoding the amino acid sequence of SEQ ID NO: 2. In one embodiment, the nucleic acid has the nucleotide sequence of SEQ ID NO: 1 (Figure

1). In another embodiment, the nucleic acid has a nucleotide sequence that is a degenerate variant of SEQ ID NO: 1. In yet another embodiment, the invention provides an isolated nucleic acid that hybridizes under stringent conditions to a nucleic acid having the nucleotide sequence of SEQ ID NO: 1, and preferably encodes a protein having the same function as a protein having the amino acid sequence of SEQ ID NO: 2.

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The isolated nucleic acid encoding hNaIII18 can be a part of a recombinant vector, e.g., for cloning, expression, and/or expansion. An expression vector comprises the nucleic acid encoding hNaIII18 operably associated with an expression control sequence. The invention further provides host cells containing such a vector, and methods for producing the hNaIII18 subunit polypeptide using such host cells.

In addition, the invention provides an isolated nucleic acid oligonucleotide, such as a primer or probe, of at least 10 bases, more particularly of at least 20, and more particularly of at least 30 bases, which oligonucleotide has a nucleotide sequence identical to a corresponding nucleotide sequence of the same number of contiguous bases in SEQ ID NO: 1, or its complement, which nucleotide sequence is unique and specific to the nucleotide sequence of SEQ ID NO: 1, and/or different from corresponding oligonucleotide sequences encoding known sodium channel subunits. The invention also provides an antibody that preferentiallyh binds the hNaIII18 subunit protein of the invention compared to other known sodium channel subunits.

The present invention further provides a method for detecting expression of hNaIII18 in a cell or sample derived from a cell, which method comprises: (i) detecting mRNA encoding hNaIII18 in a cell or in a sample derived from a cell suspected of expressing hNaIII18; or (ii) detecting hNaIII18 protein in a cell or in a sample derived from a cell with an antibody of the invention.

The present invention further provides an assay system for identifying modulators of hNaIII18 subunit containing sodium channels. The assay system comprises at least one cell genetically engineered to express or overexpress hNaIII18 as part of a functional sodium channel, which can be used to screen for and thereby identify modulators of a hNaIII18-containing sodium channel. In a preferred

embodiment, cells useful in conducting the assay are mammalian cells useful in such screening methods including, *e.g.*, human embryonic kidney cells such as HEK293 cells, or cells such as *Xenopus* cells

#### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 shows the cDNA sequence of hNaIII18 of the present invention.

Figure 2 shows the amino acid sequence of hNaIII18 of the present invention.

Figure 3 shows the cDNA sequence of human SCN3A of Clare et al. (supra) (GenBank Accession No. AJ251507).

Figure 4 shows the amino sequence of human SCN3A of Clare et al. (supra) (GenBank Accession No. AJ251507).

Figure 5 shows the cDNA of a human sodium channel  $\alpha$ -subunit variant by Jeong et al. (GenBank Accession No. AF225987).

Figure 6 shows the amino acid sequence a human sodium channel  $\alpha$ -subunit variant by Jeong et al. (GenBank Accession No. AF225987).

Figure 7 shows a cDNA alignment of the hNaIII18 of the present invention, with that of the human SCN3A of Clare et al. (*supra*), and that of Jeong et al. (*supra*)

Figure 8 shows the amino acid alignment of the hNaIII18 of the present invention, with that of the human SCN3A of Clare et al. (supra), and that of Jeong et al. (supra)

Figure 9A-D shows results of electrophysiology of hNaIII18-transfected HEK293 cells. Figure 9A demonstrates the activation threshold voltage; Figure 9B, the steady state V ½ inactivation voltage; Figure 9C, the recovery time after inactivation; and Figure 9D, the inactivation kinetics.

## **DETAILED DESCRIPTION OF THE INVENTION**

The present invention is based, in part, on the discovery of a splice variant of the human NaIII channel  $\alpha$  subunit. The human NaIII  $\alpha$  subunit isoform, designated herein as "hNaIII18", was cloned by RT-PCR from human embryonic

brain total RNA (Clontech, Palo Alto, CA), using human NaIII specific primers. Primers were designed from a sequence identified by searching the NCBI Human Genome database, using the human NaIII mRNA sequence (GenBank accession no. AJ251507) using reverse-transcriptase PCR (RT-PCR). PCR fragments were cloned into the mammalian expression vector and the complete DNA sequence was determined.

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The hNaIII18 sequence of the invention contains an additional 147 nucleotides that do not appear in the human NaIII cDNA mentioned above (SEQ ID NO: 3). Splicing in this region (nucleotides +9 to +96) had been described for the rat NaIII sodium channel, but not for the human NaIII channel when this work was initiated. The nucleotide sequence of Jeong et al. 2000, *supra*, also containing the 147 nucleotide insert and encoding an amino acid sequence similar to that of SEQ ID NO: 2, was deposited in GenBank (Accession No. AF225987, SEQ ID NO: 5), and is described in International PCT publication WO 01/96552 (in Japanese). The novel sequence (SEQ ID NO: 1) presented herein differs from that of SEQ ID NO:5 by 37 nucleotides out of 6093 aligned. None of the differences are found within the 147-nucleotide insertion. The amino acid sequence presented herein in SEQ ID NO: 2, differs from the SEQ ID NO:5 amino acid sequence by 12 amino acids out of 2000, with none of the differences being found in the region containing the 49 amino acid insert.

Transient transfection of the novel splice variant of the invention (SEQ ID NO: 1) results in expression of functional sodium channels in mammalian cells (cell line HEK293). Stable transfection and expression of the hNaIII18 also was achieved in HEK293 cells.

Protein expression was confirmed in the stably transfected HEK293 cells by immunocytochemistry and Western blotting. A protein having a size of about 220 kD protein, corresponding to the expected molecular weight of hNaIII18 was identified. Functional hNaIII18 activity was confirmed by electrophysiology.

Thus, the present invention advantageously provides hNaIII18 protein, including fragments and derivatives thereof; hNaIII18-encoding nucleic acids, and portions thereof including oligonucleotide primers and probes surrounding and within the region containing the 147 nucleotide insert, and hNaIII18 regulatory sequences;

hNaIII18-specific antibodies; and related methods of using these materials to detect the presence of hNaIII18 proteins or nucleic acids.

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The present invention also provides an assay method for screening to identify selective modulators of hNaIII18-containing sodium ion channel activity. The method involves detecting whether a test compound increases or decreases the activity of the sodium channel, as determined, e.g., by measuring current phase (electrophysiology) and ion selectivity. The assay method is preferably conducted using at least one host cell that expresses or overexpresses a functional sodium channel comprising hNaIII18, or a membrane preparation prepared therefrom. In one embodiment, the test compound inhibits (antagonizes) the activity of the sodium channel. In another embodiment, the test compound potentiates (agonizes) the activity of the sodium channel. The test system preferably involves the use of an appropriate cell culture medium to permit cell growth and viability, as well as tissue culture plates or arrays containing the host cells in the cell culture medium. In specific embodiments, host cells are mammalian cell lines such as, e.g., the HEK293 cell line, although appropriate cells from other organisms, such as, e.g., Xenopus cells, can alternatively be utilized.

The specification and figures include the following nucleotide or amino acid sequences: hNaIII18 polynucleotide (SEQ ID NO:1); hNaIII18 amino acid sequence (SEQ ID NO:2); SCN3A nucleotide sequence (SEQ ID NO:3; Clare et al., supra; GenBank AJ251507); SCN3A amino acid sequence (SEQ ID NO:4; Clare et al., supra; GenBank AJ201507); SCN3A splice variant nucleotide sequence (SEQ ID NO:5; Jeong et al., supra; GenBank AF225987); SCN3A splice variant amino acid sequence (SEQ ID NO:6; Jeong et al., supra; GenBank AF225987); forward primer utilized in Example 1 (SEQ ID NO:7); and reverse primer utilized in Example 1 (SEQ ID NO:8).

## **General Definitions**

The following definitions are provided for clarity and illustrative purposes only, and are not intended to limit the scope of the invention.

As used herein, the term "isolated" means that the referenced material is removed from the environment in which it is normally found. Thus, an isolated

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biological material can be free of cellular components, i.e., components of the cells in which the material is found or produced in nature. In the case of nucleic acid molecules, an isolated nucleic acid includes a PCR product, an mRNA, a cDNA, or a restriction fragment. In another embodiment, an isolated nucleic acid is preferably excised from the chromosome in which it may be found, and more preferably is no longer joined to non-regulatory, non-coding regions, or to other genes, located upstream or downstream of the gene contained by the isolated nucleic acid molecule when found in the chromosome. In yet another embodiment, the isolated nucleic acid lacks one or more naturally occurring introns. Isolated nucleic acid molecules include sequences inserted into plasmids, cosmids, artificial chromosomes, phages and the like. Thus, in a specific embodiment, a recombinant nucleic acid is an isolated nucleic acid. An isolated protein may be associated with other proteins or nucleic acids, or both, with which it associates in the cell, or with cellular membranes if it is a membrane-associated protein. A protein expressed from a vector in a cell, particularly a cell in which the protein is normally not expressed, is also a regarded as isolated. An isolated organelle, cell, or tissue is removed from the anatomical site in which it is found in a cell or an organism. An isolated material may be, but need not be, purified. As used herein to refer to nucleic acids, the term "isolated" does not encompass man-made genomic or cDNA libraries.

The term "purified" as used herein refers to material that has been isolated under conditions that reduce or eliminate the presence of unrelated materials, *i.e.*, contaminants, including native materials from which the material is obtained. For example, a purified protein is preferably substantially free of other proteins or nucleic acids with which it is associated in a cell; a purified nucleic acid molecule is preferably substantially free of proteins or other unrelated nucleic acid molecules with which it can be found within a cell. As used herein, the term "substantially free" is used operationally, in the context of analytical testing of the material. Preferably, purified material substantially free of contaminants. Purity can be evaluated by chromatography, gel electrophoresis, immunoassay, composition analysis, biological assay, and other methods known in the art.

Methods for purification are well-known in the art. For example, nucleic acids can be purified by precipitation, chromatography (including preparative

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solid phase chromatography, oligonucleotide hybridization, and triple helix chromatography), ultracentrifugation, and other means. Polypeptides and proteins can be purified by various methods including, without limitation, preparative disc-gel electrophoresis, isoelectric focusing, HPLC, reversed-phase HPLC, gel filtration, ion exchange and partition chromatography, precipitation and salting-out chromatography, extraction, and countercurrent distribution. For some purposes, it is preferable to produce the protein in a recombinant system so that it contains an additional sequence tag that facilitates purification, such as, but not limited to, a polyhistidine sequence (His®-tag; Novagen, Madison, WI), or a sequence that specifically binds to an antibody, such as the FLAG® tag (Sigma, St. Louis, MO), HA-tag (Roche Diagnostics, Indianapolis, IN), or that can be column-purified such as via the use of glutathione-S-transferase (GST). The polypeptide can then be purified from a crude lysate of the host cell by chromatography on an appropriate solid-phase matrix. Alternatively, antibodies produced against the protein or against peptides derived therefrom can be used as purification reagents. Cells can be purified by various techniques, including centrifugation, matrix separation (e.g., nylon wool separation), panning and other immunoselection techniques, depletion (e.g., complement depletion of contaminating cells), and cell sorting (e.g., fluorescence activated cell sorting (FACS)). Other purification methods are possible. A purified material may contain less than about 50%, preferably less than about 75%, and most preferably less than about 90%, by weight of the cellular components with which it was originally associated. The "substantially pure" indicates the highest degree of purity that can be achieved using conventional purification techniques known in the art.

In a specific embodiment, the term "about" or "approximately" means plus or minus 10% of the stated numerical value or range.

As use herein, the term "ion channel" refers to a transmembrane pore that presents a hydrophilic channel for ions to cross a lipid bilayer down their electrochemical gradients. In a preferred embodiment, the ion channel is a voltage-gated sodium ion channel. A "sodium channel" is an ion channel that is selective for sodium ions.

A "sample" as used herein refers to a biological material that can be obtained and tested for the presence or expression of: (i) an hNaIII18 subunit-containing ion channel; or (ii) an hNaIII18 subunit protein; or (iii) an hNaIII18 subunit-encoding nucleic acid. Such samples can be obtained from animal, preferably mammalian, and more preferably human subjects, and include tissue samples, especially CNS or PNS tissues, as well as cell cultures derived from such tissues. Alternatively, such samples can comprise cells genetically engineered to express or overexpress an hNaIII18 subunit-containing ion channel or an hNaIII18 subunit protein. Such cells are preferably eukaryotic, but may alternatively be prokaryotic cells. Eukaryotic cells are preferably mammalian cells, but may alternatively be *Xenopus* cells.

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Non-human animals include, without limitation, laboratory animals such as mice, rats, rabbits, hamsters, guinea pigs, etc.; domestic animals such as dogs and cats; and farm animals such as sheep, goats, pigs, horses, and cows.

The term "modulator" refers to a compound that binds to an ion channel comprising the hNaIII18 subunit protein of the invention and differentially affects the activity of the ion channel in response to a stimulus that normally activates the function of that ion channel when compared to the activity of the ion channel not contacted with the compound. Ion channel activity can be measured, *e.g.*, using electrophysiological techniques, or according to other known methods in the art. In a preferred embodiment, the ion channel is a sodium channel.

The terms "inhibitor" and antagonist refer to a compound that binds to the ion channel comprising hNaIII18, and blocks, inhibits, impedes or reduces the activity of that ion channel.

An "agonist" is defined as a compound that binds to the ion channel comprising hNaIII18, and promotes, enhances, stimulates or potentiates the normal biological function of the sodium channel. A "partial agonist" binds as to the ion channel or a subunit thereof, as does a full agonist, but promotes only partial function.

As used herein the term "transfected cell" or "transformed cell" refers to a host cell that has been genetically engineered to express or overexpress a nucleic acid encoding a hNaIII18 subunit, preferably in combination with one or more  $\beta$  subunits such as, *e.g.*,  $\beta$ -subunits 1-3 as described in GenBank Accession Nos.

U87445, AF007783, AH005825, AF007783, AF04948, L10338 and L16242, among others. Any cell can be used, preferably a eukaryotic cell, and more preferably a vertebrate cells, preferably a mammalian cell, or a *Xenopus* cell. Such cells additionally can be genetically engineered to coexpress or overexpress a different sodium channel subunit. Such genetically engineered cells include those cells into which one or more heterologous hNaIII18-encoding nucleic acids have been introduced and are expressed or overexpressed. Such genetically engineered cells also include those cells engineered to express or overexpress one or more endogenous hNaIII18 subunits, for example, by gene activation technology.

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Such cells are particularly suitable to conduct an assay to screen for compounds that modulate the function of the hNaIII18 subunit-containing sodium channel in response to an appropriate stimulus (e.g., TTX). An "assay method" typically makes use of one or more such cells, e.g., in a microwell plate or some other culture system. The effects of a test compound can be determined on a single cell or on a collection of cells sufficient to allow measurement of ionic current, activation threshold, or ionic permeability characteristics of the hNaIII18 subunit-containing sodium channels. For example, single cells can be tested, e.g., by use of patch clamp or other appropriate electrophysiological techniques.

A "test compound" or "candidate compound" is any molecule that can be tested for its ability to bind to the hNaIII18 subunit-containing sodium channel, or to a subunit thereof, and preferably modulate on the activity of the hNaIII18 subunit-containing sodium channel. A compound that binds and modulates a hNaIII18 subunit-containing sodium channel is a "lead compound" suitable for further testing and development.

The term "ligand" can alternatively be used to refer to any compound or peptide or polypeptide that binds to and modulates the activity of a hNaIII18 subunit, or a sodium channel comprising hNAIII18.

The term "pain disorder" includes chronic pain, defined as pain lasting longer than one month (Bonica, Semin Anesth 1986, 5:82-99), and is characterized by unrelenting persistent pain that is not amenable to routine pain control methods. The term "pain disorder" also includes neuropathic pain and nociceptive pain.

"Chronic pain" can be defined as pain lasting longer than one month (Bonica, Semin Anesth 1986, 5:82-99), and is characterized by unrelenting persistent pain that is not amenable to routine pain control methods. Chronic pain includes, but is not limited to, inflammatory pain, postoperative pain, cancer pain, osteoarthritis pain associated with metastatic cancer, trigeminal neuralgia, acute herpetic and postherpetic neuralgia, diabethic neuropathy, causalgia, brachial plexus avulsion, occipital neuralgia, reflex sympathetic dystrophy, fibromyalgia, gout, phantom limb pain, burn pain, pain associated with spinal cord injury, multiple sclerosis, reflex sympathetic dystrophy and lower back pain and other forms of neuralgia, neuropathic, and idiopathic pain syndromes.

"Neuropathic pain" can be caused by injury or infection of peripheral sensory nerves. It includes, but is not limited to pain from peripheral nerve trauma, herpes virus infection, diabetes mellitus, causalgia, plexus avulsion, neuroma, limb amputation, and vasculitis. Neuropathic pain is also caused by nerve damage from chronic alcoholism, human immunodeficiency virus infection, hypothyroidism, uremia, or vitamin deficiences. Neuropathic pain includes but is not limited to pain caused by nerve injury such as, for example, the pain from which diabetics suffer.

Chronic and neuropathic types of pain generally arises from injury to the peripheral or central nervous tissue.

"Nociceptive pain" is due to activation of pain-sensitive nerve fibers, either somatic or visceral. Nociceptive pain generally results as a response to direct tissue damage. The initial trauma causes the release of several chemicals including bradykinin, serotonin, substance P, histamine, and prostaglandin. When somatic nerves are involved, the pain is typically experienced as aching or pressure-like.

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#### Molecular Biology Definitions

In accordance with the present invention there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. See, e.g., Sambrook, Fritsch & Maniatis, Molecular Cloning: A Laboratory Manual, Second Edition (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (herein "Sambrook et al., 1989"); DNA Cloning: A Practical Approach, Volumes I and II (D.N. Glover ed. 1985);

Oligonucleotide Synthesis (M.J. Gait ed. 1984); Nucleic Acid Hybridization [B.D. Hames & S.J. Higgins eds. (1985)]; Transcription And Translation [B.D. Hames & S.J. Higgins, eds. (1984)]; Animal Cell Culture [R.I. Freshney, ed. (1986)]; Immobilized Cells And Enzymes [IRL Press, (1986)]; B.Perbal, A Practical Guide To Molecular Cloning (1984); F.M. Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, Inc. (1994).

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"Amplification" of DNA as used herein denotes the use of exponential amplification, techniques such as polymerase chain reaction (PCR), and non-exponential amplification, such as linked linear amplification, to increase the concentration of a particular DNA sequence within a mixture of DNA sequences. For a description of PCR see Saiki et al., Science 1988, 239:487. For a description of linked linear amplification, see U.S. Patent Nos. 6,335,184 and 6,027,923 and Reyes et al. Clinical Chemistry 2001; 47: 131-40; Wu et al. Genomics 1989; 4: 560-569.

As used herein, "sequence-specific oligonucleotides" refers to related sets of oligonucleotides that can be used to detect allelic variations or mutations in the hNaIII18 gene, or can be used for amplification of an hNAIII18 encoding-nucleic acid.

The nucleic acid molecules (polynucleotides) described herein may be flanked by natural regulatory (expression control) sequences, or may be associated with heterologous sequences, including promoters, internal ribosome entry sites (IRES) and other ribosome binding site sequences, enhancers, response elements, suppressors, signal sequences, polyadenylation sequences, introns, 5'- and 3'- non-coding regions, and the like. The nucleic acid molecules may also be modified by many means known in the art. Non-limiting examples of such modifications include methylation, "caps", substitution of one or more of the naturally occurring nucleotides with an analog, and internucleotide modifications such as, for example, replacement with uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoroamidates, carbamates, etc.) and with charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.). Polynucleotides may contain one or more additional covalently linked moieties, such as, for example, proteins (e.g., nucleases, toxins, antibodies, signal peptides, poly-L-lysine, etc.), intercalators (e.g., acridine, psoralen, etc.), chelators (e.g., metals, radioactive metals, iron, oxidative

metals, etc.), and alkylators. The polynucleotides may be derivatized by formation of a methyl or ethyl phosphotriester or an alkyl phosphoramidate linkage. Furthermore, the polynucleotides herein may also be modified with a label capable of providing a detectable signal, either directly or indirectly. Exemplary labels include radioisotopes, fluorescent molecules, biotin, and the like.

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A "coding sequence" or a sequence "encoding" an expression product, such as an RNA, polypeptide, protein, or enzyme, is a nucleotide sequence that, when expressed, results in the production of that RNA or polypeptide, *i.e.*, the nucleotide sequence encodes an amino acid sequence for that polypeptide. A coding sequence or "open reading frame (ORF)" for a polypeptide will typically include a start codon (usually ATG) and a stop codon.

The term "gene", also called a "structural gene" refers to a basic unit of hereditary material. Specifically a gene is an ordered sequence of DNA nucleotide bases that encodes one polypeptide chain (via mRNA). The gene includes regions preceding and following the coding region (such as promoter sequences, a 5'-untranslated region, and a 3'-untranslated region, which affect, for example, the conditions under which the gene is expressed) as well as (in eukaryotes) intervening sequences (introns) between individual coding segments (exons).

A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary to initiate transcription at levels detectable above background. Within the promoter sequence will be found a transcription initiation site (conveniently defined for example, by mapping with nuclease S1), as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase. The present invention includes the hNaIII18 gene promoter found in the genome, which can be operatively associated with a hNaIII18 coding sequence with a heterologous coding sequence.

The term "host cell" means any cell of any organism that is selected, modified, transformed, grown, or used or manipulated in any way, for the production

of a substance by the cell, for example, the expression by the cell of a gene, a DNA or RNA sequence, or a polypeptide. Host cells can further be used for screening or other assays, as described *infra*.

A coding sequence is "under the control of" or "operatively associated with" transcriptional and translational control sequences in a cell when such control sequences operate to effect RNA polymerase transcription of the coding sequence into mRNA, which is then trans-RNA spliced (if it contains introns) and translated, in the case of mRNA, into the protein encoded by the coding sequence.

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The terms "express" and "expression" mean allowing or causing the information in a gene or cDNA or mRNA sequence to become manifest, for example, by producing a protein by activating the cellular functions\_involved in transcription and translation of a corresponding gene, cDNA or mRNA sequence. A gene or cDNA sequence is expressed in or by a cell to form an "expression product" such as a protein. The expression product itself, *e.g.*, the resulting protein, may also be said to be "expressed" by the cell. An expression product can be characterized as intracellular, extracellular, transmembrane, or secreted depending on the particular product. The hNaIII18 subunit protein of the invention is typically expressed as a transmembrane protein with intracellular and extracellular domains.

The term "transfection" means the introduction of a "foreign" (i.e., extrinsic or extracellular) gene, DNA or RNA sequence into a host cell so that the host cell will express the introduced gene or sequence to produce a desired substance, typically a protein encoded by the introduced gene or sequence. The introduced gene or sequence may also be called a "cloned" or "foreign" or "heterologous" gene or sequence, and may include regulatory or control sequences, such as start, stop, promoter, signal, secretion, or other sequences used by a cell's genetic machinery. The gene or sequence may include non-functional sequences or sequences with no known function.

The term "transformation" refers to the process by which DNA is introduced from the surrounding medium into a prokaryotic host cell.

The term "transduction" refers to the introduction of DNA into a prokaryotic host cell via a bacterial virus, or bacteriophage.

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A prokaryotic or eukaryotic host cell that receives and expresses introduced DNA or RNA has been "transformed" and is a "transformant" or a "clone." The DNA or RNA introduced into a host cell can come from any source, including cells of the same genus or species as the host cell, or cells of a different genus or species, or synthetic sequences.

The transformed cells of the invention are particularly suitable for an assay system for the detection of compounds that modulate the function of hNaIII18 subunit-containing sodium channels in response to activation, *e.g.*, in response to exposure TTX. An "assay method" makes use of one or more such cells, *e.g.*, in a microwell plate or some other culture or assay system to permit evaluation of the effects of a test compound on the cell(s), *e.g.*, by measuring ionic current or activation threshold characteristics of the hNaIII18 subunit-containing sodium channel.

The term "recombinantly engineered cell" refers to any prokaryotic or eukaryotic cell that has been manipulated to express or overexpress the hNaIII18 subunit by any appropriate method, including transfection, transformation or transduction. This term also includes endogenous activation of a hNaIII18 gene in a cell that does not normally express hNaIII18 or that expresses the protein at a suboptimal level.

The terms "vector", "cloning vector" and "expression vector" mean the vehicle by which a DNA or RNA sequence (e.g., a foreign gene) can be introduced into a host cell, so as to transform the host and promote expression (e.g., transcription and translation) of the introduced sequence. Vectors include plasmids, cosmids, phages, viruses, etc.; they are discussed in greater detail below.

Vectors typically comprise the DNA of a transmissible agent, into which foreign DNA is inserted. A common way to insert one segment of DNA into another segment of DNA involves the use of restriction enzymes to cleave DNA at specific restriction sites. A "cassette" refers to a DNA coding sequence or segment of DNA that codes for an expression product that can be inserted into a vector at defined restriction sites. The cassette restriction sites are designed to ensure insertion of the cassette in the proper reading frame. Generally, foreign DNA is inserted at one or more restriction sites of the vector DNA, and then is carried by the vector into a host cell along with the transmissible vector DNA. A segment or sequence of DNA

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having inserted or added DNA, such as an expression vector, can also be called a "DNA construct." A common type of vector is a plasmid. A plasmid vector often contains coding DNA and promoter DNA and has one or more restriction sites suitable for inserting foreign DNA. Coding DNA is a DNA sequence that encodes a particular amino acid sequence for a particular protein. Promoter DNA is a DNA sequence that initiates, regulates, or otherwise mediates or controls the expression of the coding DNA. Promoter DNA and coding DNA may be from the same gene or from different genes, and may be from the same or different organisms. A large number of vectors, including plasmid and fungal vectors, have been described for replication and/or expression in a variety of eukaryotic and prokaryotic hosts. Nonlimiting examples include pKK plasmids (Clonetech), pUC plasmids, pET plasmids (Novagen, Inc., Madison, WI), pRSET or pREP plasmids (Invitrogen, San Diego, CA), or pMAL plasmids (New England Biolabs, Beverly, MA), and many appropriate host cells. Recombinant cloning vectors will often include one or more replication systems for cloning or expression, one or more markers for selection in the host, e.g., antibiotic resistance, and one or more expression cassettes.

The term "expression system" means a host cell and compatible vector under suitable conditions, *e.g.*, for the expression of a protein coded for by foreign DNA carried by the vector and introduced to the host cell. Common expression systems include *E. coli* host cells and plasmid vectors, insect host cells and baculovirus vectors, and mammalian host cells and vectors.

The term "heterologous" refers to a combination of elements not naturally occurring. For example, heterologous DNA refers to DNA not naturally present in that cell. Alternativley, heterologous DNA refers to combinations of sequences that do not naturally occur together in that cell, *e.g.*, promoter sequences from a gene from one cell type linked to coding sequences of a gene that is not normally controlled by that promoter or expressed by another cell type. Preferably, the heterologous DNA includes a gene foreign to the cell. A heterologous expression regulatory element is such an element operatively associated with a different gene than the one it is operatively associated with in nature. In the context of the present invention, a hNaIII18 gene is heterologous to the vector DNA in which it is inserted

for cloning or expression purposes, and is heterologous to a host cell containing such a vector in which it is expressed, e.g., a HEK cell.

The terms "mutant" and "mutation" mean any detectable change in genetic material, e.g., DNA, or any process, mechanism, or result of such a change. This includes gene mutations in which the structure (e.g., DNA sequence) of a gene is altered; any gene or DNA arising from any mutation process; and any expression product (e.g., protein or enzyme) expressed by a non-silent modification of a gene or DNA sequence. The term "variant" may also be used to indicate a modified or altered gene, DNA sequence, polypeptide, cell, etc., i.e., any kind of mutant therefrom.

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"Sequence-conservative variants" or "degenerate variants" of a polynucleotide sequence are those in which a change of one or more nucleotides in a given codon position results in no alteration in the amino acid encoded at that position.

"Function-conservative variants" are those in which a given amino acid residue in a protein has been changed without substantially altering the function of the polypeptide, including, but not limited to, replacement of an amino acid with a residue having similar properties (such as, for example, polarity, hydrogen bonding potential, acidic, basic, hydrophobic, aromatic, and the like). Amino acids with similar properties are well known in the art. For example, arginine, histidine and lysine are hydrophilic-basic amino acids and may be interchangeable. Similarly, isoleucine, a hydrophobic amino acid, may be replaced with leucine, methionine or valine. Such changes are expected to have little or no effect on the apparent molecular weight, isoelectric point, or function of the protein. Amino acid residues may be varied in a protein so that the percent amino acid sequence identity between the original protein and the variant may be, for example, at least 70%, 80%, 90%, 95% or 99%, as determined according to a default alignment scheme such as by the Cluster Method, wherein similarity is based on the MEGALIGN algorithm, or BLAST. A "functionconservative variant" of the present invention includes those polypeptides having the above-described amino acid sequence identities, and having the same or substantially similar functions as the native or parent hNaIII18 subunit protein of the invention

As used herein, the term "homologous" refers to the relationship between proteins that possess a "common evolutionary origin," including proteins

from superfamilies (e.g., the immunoglobulin superfamily) and homologous proteins from different species (e.g., myosin light chain, etc.) (Reeck et al., Cell 1987, 50:667). Such proteins (and their encoding genes) have sequence homology, as reflected by their sequence similarity or sequence identity, whether in terms of percent similarity or the presence of specific residues or motifs at conserved positions.

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Accordingly, the term "sequence similarity" or "sequence identity" refers to the degree of identity or correspondence between nucleic acid or amino acid sequences of proteins that may or may not share a common evolutionary origin (see Reeck et al., *supra*). However, in common usage and in the instant application, the term "homologous," when modified with an adverb such as "highly," may refer to sequence similarity and may or may not relate to a common evolutionary origin.

In a specific embodiment, two DNA sequences are "substantially homologous" or "substantially similar" when at least about 80%, and most preferably at least about 90, 95% or 99% of the nucleotides match over the defined length of the DNA sequences, as determined by sequence comparison algorithms, such as BLAST, FASTA, DNA Strider, etc. An example of such a sequence is an allelic or species variant of the specific hNaIII18 gene of the invention. Sequences that are substantially homologous can be identified by comparing the sequences using standard software available in sequence data banks, or in a Southern hybridization experiment under, for example, stringent conditions as defined for that particular system.

Similarly, in a particular embodiment, two amino acid sequences are "substantially homologous" or "substantially similar" when greater than 80%, 90%, 95% or 99% of the amino acids are identical. Preferably, the similar or homologous sequences are identified by alignment using, for example, the GCG (Genetics Computer Group, Program Manual for the GCG Package, Version 7, Madison, Wisconsin) pileup program, or any of the programs described above (BLAST, FASTA, etc.).

A nucleic acid molecule is "hybridizable" to another nucleic acid molecule, such as a cDNA, genomic DNA, or RNA, when a single stranded form of the nucleic acid molecule can anneal to the other nucleic acid molecule or its complement under the appropriate conditions of temperature and solution ionic

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strength (see Sambrook et al., supra). The conditions of temperature and ionic strength determine the "stringency" of the hybridization. For preliminary screening for homologous nucleic acids, low stringency hybridization conditions, using a Tm (melting temperature) in the range of about 55 °C with low salt and/or denaturant concentrations, can be used, e.g., 5x SSC, 0.1% SDS, 0.25% milk, and no formamide; or 30% formamide, 5x SSC, 0.5% SDS. Moderate stringency hybridization conditions correspond to use of a higher Tm, and higher concentrations of salt and/or denaturants, e.g., 40% formamide, with 5x or 6x SSC. High stringency hybridization conditions correspond to the highest Tm and concentrations of salt/and/or denaturants, e.g., 68°C, 50% formamide, 5x or 6x SSC. SSC is a 0.15M NaCl, 0.015M Na-citrate buffer. Hybridization requires that the two nucleic acids contain complementary sequences, although depending on the stringency of the hybridization, mismatches between bases are possible. The appropriate stringency for hybridizing nucleic acids depends on the length of the nucleic acids and the degree of complementation, as known in the art. The greater the degree of similarity or homology between two nucleotide sequences, the higher the value of Tm for hybrids of nucleic acids having those sequences. The relative stability (corresponding to higher Tm) of nucleic acid hybridizations decreases in the following order: RNA:RNA, DNA:RNA, DNA:DNA. For hybrids of greater than 100 nucleotides in length, equations for calculating Tm have been derived (see Sambrook et al. 1989, supra, 9.50-9.51). For hybridization with shorter nucleic acids, i.e., oligonucleotides, the position of mismatches becomes more important, and the length of the oligonucleotide determines its specificity (see Sambrook et al., supra, 11.7-11.8). A minimum length for a hybridizable nucleic acid is at least about 10 nucleotides; preferably at least about 15 nucleotides; and more preferably at least about 20 nucleotides.

In a specific embodiment, the term "standard hybridization conditions" refers to a Tm of 55°C, and utilizes conditions as set forth above. In a preferred embodiment, the Tm is about 60°C; in a more preferred embodiment, the Tm is about 65°C. In a specific embodiment, "high stringency" refers to hybridization and/or washing conditions at 68°C, in 0.2 x SSC, at 42°C in 50% formamide, 4x SSC, or under conditions that afford levels of hybridization equivalent to those observed under either of these two conditions.

As used herein, the term "oligonucleotide" refers to a nucleic acid, generally of at least 10, preferably at least 15, and more preferably at least 20 nucleotides, preferably no more than 100 nucleotides, that is hybridizable to a genomic DNA molecule, a cDNA molecule, or an mRNA molecule, or other nucleic acid of interest. Oligonucleotides can be labeled, *e.g.*, with  $\gamma^{32}$ P-nucleotides or nucleotides to which a label, such as biotin, has been covalently conjugated. In one embodiment, a labeled oligonucleotide can be used as a probe to detect the presence of a nucleic acid. In another embodiment, oligonucleotides (one or both of which may be labeled) can be used as PCR primers, either for cloning a full length nucleic acid or a fragment of a nucleic acid encoding the hNaIII18 subunit, or to detect the presence of nucleic acids encoding hNaIII18. In a further embodiment, an oligonucleotide of the invention can form a triple helix with a hNaIII18-encoding DNA molecule. Generally, oligonucleotides are prepared synthetically, preferably on a nucleic acid synthesizer. Accordingly, oligonucleotides can be prepared with non-naturally occurring phosphoester analog bonds, such as thioester bonds, etc.

The present invention also provides antisense nucleic acids, which may be used to inhibit expression of the hNaIII18 subunit protein of the invention.

Inhibition of hNaIII18 expression may be desired when upregulation of hNaIII18 expression or excessive activation of an hNaIII18-containing ion channel induces or otherwise contributes to an increase in pain or a pain disorder in a subject.

An "antisense nucleic acid" is a single stranded nucleic acid molecule, which may be DNA, RNA, a DNA-RNA chimera, or derivatives thereof, which, on hybridizing under cytoplasmic conditions with complementary bases in an RNA or DNA molecule, inhibits the expression or translation of the encoded gene. If the RNA is an mRNA transcript, the antisense nucleic acid is a counter-transcript or mRNA-interfering complementary nucleic acid. As presently used, "antisense" broadly includes RNA-RNA interactions, RNA-DNA interactions, and RNase-H mediated arrest. Antisense nucleic acid molecules can be encoded by a recombinant gene for expression in a cell (e.g., U.S. Patent No. 5,814,500; U.S. Patent No. 5,811,234), or alternatively they can be prepared synthetically (see, e.g., U.S. Patent No. 5,780,607).

In addition to antisense sequences, the present invention also provides ribozymes useful to inhibit hNaIII18 expression. Ribozyme technology is described further in Intracellular Ribozyme Applications: Principals and Protocols, Ed. Rossi and Couture, 1999, Horizon Scientific Press

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#### hNaIII18 Nucleic Acids

A polynucleotide molecule encoding hNaIII18, whether genomic DNA or cDNA, can be isolated from any source, particularly from a human cDNA or genomic library. Methods for obtaining specific polynucleotide molecules gene are well known in the art, as described above (see, e.g., Sambrook et al., 1989, supra). The DNA may be obtained by standard procedures known in the art from cloned DNA (e.g., a DNA "library"), and preferably is obtained from a cDNA library prepared from tissues with high level expression of the encoded protein, by chemical synthesis, by cDNA cloning, or by the cloning of genomic DNA, or fragments thereof, purified from the desired cell (See, for example, Sambrook et al., 1989, supra; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II). Clones derived from genomic DNA may contain regulatory and intron DNA regions in addition to coding regions. Clones derived from cDNA will not contain intron sequences. Whatever the source, the polynucleotide molecule should be cloned into a vector suitable for its propagation. Identification of a specific DNA fragment containing the desired hNaIII18-encoding sequence may be accomplished in a number of ways. For example, a portion of a hNaIII18 encoding polynucleotide molecule exemplified infra can be purified and labeled to prepare a labeled probe, and the generated DNA library may be screened by nucleic acid hybridization to the labeled probe (Benton and Davis, Science 1977, 196:180; Grunstein and Hogness, Proc. Natl. Acad. Sci. U.S.A. 1975, 72:3961). Those DNA fragments with substantial homology to the probe, such as an allelic variant from another individual, will hybridize. In a specific embodiment, highest stringency hybridization conditions are used to identify a homologous hNaIII18 gene.

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Further selection can be carried out on the basis of the properties of the gene, e.g., if the gene encodes a protein product having the same physicochemical profile (i.e., isoelectric, electrophoretic, electrophysiological, amino acid composition,

partial or complete amino acid sequence, antibody binding activity, or ligand binding profile) of the hNaIII18 subunit protein disclosed herein. Thus, the presence of the nucleic acid may be detected by assays based on the physical, chemical, immunological, or functional properties of its expressed product.

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Other DNA sequences which encode substantially the same amino acid sequence as a hNaIII18 gene may be used in the practice of the present invention. These include but are not limited to allelic variants, species variants, sequence conservative variants, and function conservative variants.

Amino acid substitutions may also be introduced to substitute an amino acid with a particularly preferable property. For example, a Cys may be introduced at a potential site for disulfide bridges with another Cys.

Polynucleotide molecules encoding the hNaIII18 subunit, and the encodied polypeptide, derivatives and analogs thereof, can be produced by various methods known in the art. The manipulations which result in their production can occur at the gene or protein level. For example, the cloned hNaIII18 gene or cDNA sequence can be modified by any of numerous strategies known in the art (Sambrook et al., 1989, supra). The sequence can be cleaved at appropriate sites with restriction endonuclease(s), followed by further enzymatic modification if desired, isolated, and ligated in vitro. In the production of the polynucleotide molecule encoding a derivative or analog of hNaIII18, care should be taken to ensure that the modified polynucleotide sequence remains within the same translational reading frame as the hNaIII18 gene, uninterrupted by premature translational stop signals.

Additionally, the encoding nucleic acid sequence can be mutated *in vitro* or *in vivo* to create and/or destroy translation, initiation, and/or termination sequences, or to create variations in coding regions and/or form new restriction endonuclease sites or destroy preexisting ones, to facilitate further *in vitro* modification. Such modifications can be made to introduce restriction sites and facilitate cloning the polynucleotide molecule into an expression vector. Any technique for mutagenesis known in the art can be used, including but not limited to, *in vitro* site-directed mutagenesis (Hutchinson, C., *et al.*, J. Biol. Chem.1978; 253:6551; Zoller and Smith, DNA 1984; 3:479-488; Oliphant *et al.*, Gene 1986; 44:177; Hutchinson *et al.*, Proc. Natl. Acad. Sci. U.S.A.1986; 83:710), use of TAB

linkers (Pharmacia), etc. PCR techniques are preferred for site directed mutagenesis (see Higuchi, 1989, "Using PCR to Engineer DNA", in PCR Technology: Principles and Applications for DNA Amplification, H. Erlich, ed., Stockton Press, Chapter 6, pp. 61-70).

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The identified and isolated polynucleotide molecule can then be inserted into an appropriate cloning vector. A large number of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. Examples of vectors include, but are not limited to, E. coli, bacteriophages such as lambda derivatives, or plasmids such as Bluescript, pBR322 derivatives or pUC plasmid derivatives, e.g., pGEX vectors, pmal-c, pFLAG, etc. The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector that has complementary cohesive termini. However, if the complementary restriction sites used to fragment the DNA are not present in the cloning vector, the ends of the DNA molecules may be enzymatically modified. Alternatively, any restriction site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically synthesized oligonucleotides encoding restriction endonuclease recognition sequences. In addition, simple PCR or overlapping PCR may be used to insert a fragment into a cloning vector.

Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated. Preferably, the cloned gene is contained on a shuttle vector plasmid, which provides for propagation in a cloning cell, e.g., E. coli, and facile purification for subsequent insertion into an appropriate expression cell line, if such is desired. For example, a shuttle vector, which is a vector that can replicate in more than one type of organism, can be prepared for replication in both E. coli and Saccharomyces cerevisiae by linking sequences from an E. coli plasmid with sequences from the yeast  $2\Phi$  plasmid.

In a preferred embodiment of the invention, the hNaIII18 sodium channel is cloned using a strategy designed to minimize mutations during cDNA

preparation, RT-PCR amplification, and growth in bacteria. This strategy is described in detail *infra*, in Example 1. The main points are summarized as follows:

First, as an alternative to conventional reverse transcriptases, which function optimally at temperatures of between 37 °C and 43 °C, this method employs an avian RNase (-) reverse transcriptase that functions optimally at temperatures between 50-65 °C. The higher temperature serves to decrease secondary structure of the RNA to produce higher cDNA yield.

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Second, for amplification of the cDNA, an enzyme mixture comprising the conventional thermostable Taq polymerase and Pwo polymerase is used. This mixture is optimized to produce very large PCR products with low error frequency, thus decreasing the mutation frequency.

Third, the number of cycles of amplification is decreased to about 28, as opposed to the typical 30-35 cycles to further reduce the possibility of mutation.

Fourth, the PCR products are electrophoresed and visualized on an agarose gel containing Crystal Violet stain, as opposed to ethidium bromide. Crystal Violet allows visualization in white light, eliminating the need for UV exposure. UV is known to induce mutations in ethidium bromide-stained DNA.

Fifth, to minimize recombination and mutation in plasmid DNA during amplification in bacteria, the PCRamplified cDNA is cloned into a low-copy number expression vector that is engineered to have limited replication cycles and contains a tetracycline-resistance gene as a selectable marker instead of an ampicillinresistance gene. Fewer replication cycles again reduces the error rate during DNA synthesis, and selection with tetracycline is less likely to induce mutations in the plasmid than is ampicillin.

Sixth, competent bacterial cells that are designed to optimize cloning of unstable inserts are selected for the transformation, and grown at a lower temperature (30-33°C versus 37°C) to decrease the growth rate and therefore, minimize the possibility of mutations. In addition, the cultures should be maintained in exponential (log) phase throughout growth, eliminating the possibility of mutations resulting from starvation, poor aeration, and accumulation of toxic metabolites.

Seventh, small tetracycline resistant colonies are chosen for evaluation rather than large ones. Human NaIII expression during growth is expected to be toxic to bacteria, thus transformed cells will yield smaller colonies.

### hNaIII18 Regulatory Nucleic Acids

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Elements of the hNaIII18 promoter can be identified by scanning the human genomic region upstream of the hNaIII18 start site, *e.g.*, by creating deletion mutants and checking for expression, or by using an algorithm. Sequences up to about 6 kilobases (kb) or more upstream from the hNaIII18 start site can contain tissue-specific regulatory elements.

The term "hNaIII18 promoter" encompasses artificial or heterologous promoters. Such promoters can be prepared by deleting non-essential intervening sequences from the upstream region of the hNaIII18 promoter, or by joining upstream regulatory elements from the hNaIII18 promoter with a heterologous minimal promoter, such as the CMV immediate early promoter.

A hNaIII18 promoter can be operably associated with a heterologous coding sequence, *e.g.*, for a reporter gene (luciferase and green fluorescent proteins are examples of reporter genes) in a construct. This construct can be used to test for conditions or reagents that normally result in expression. This construct can be used in screening assays, described below, for hNaIII18 agonists and antagonists.

hNaIII18 regulatory nucleic acids of the present invention also include non-endogenous or artificial promoter sequences or sequences that encode zinc finger proteins that may be used, e.g., in gene activation techniques, to initiate expression of hNaIII18 in cells where it is not normally expressed or to upregulate expression of the hNaIII18 subunit protein to a higher level where it would otherwise be expressed in suboptimal levels. Gene activation techniques that may be adapted to this use are described in the art, e.g., in U.S. Patent Nos. 5,968,502 and 6,214,622 to Treco et al.

## Expression of hNaIII18 Polypeptides

The primary goal for establishing a stable cell line expressing functional human sodium channels is to identify antagonists to inhibit sodium currents

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mediated by the sodium channels. DRG neurons transmit nociceptive signals from the peripheral nervous system to the central nervous system. TTX-S and TTX-R sodium channels mediate the DRG action potentials responsible for these signals. However, DRG neurons express several different isoforms of TTX-S and TTX-R currents, thereby making it difficult to determine specific interactions of antagonists with particular subtypes of sodium channels in these cells.

By generating a cell line that expresses a single sodium channel subtype, e.g., hNaIII18, alone or preferably in combination with appropriate  $\beta$ subunits, the effect of drugs on the different sodium channel isoforms can be assessed. Previously, developing stable cell lines expressing nucleic acids containing repetitive sequences, such as those contained within sodium channel genes, has been challenging. In particular, cell lines expressing functional sodium channels have been difficult to generate due to the occurrence of inactivating mutations arising in the cDNA during the cloning process (i.e., cDNA preparation, PCR amplification, and subsequent growth in bacteria). International PCT publication WO 98/38302 (Delgado et al.) describes isolation, cloning and expression of a rat TTX-S sodium channel in Xenopus oocytes. Experiments described therein demonstrate the formation of a functional TTX-S channel after injection of cRNA into Xenopus oocytes for the  $\alpha$ -subunit, alone or in combination with the  $\beta$ 1,  $\beta$ 2 or  $\beta$ 3 subunits. International PCT Publication WO 01/68681 (Aitken et al.) describes altered ion channel proteins having acquired sensitivity or refractory sensitivity to a gating agent. A rat sodium channel type II was modified by site-directed mutagenesis and PCR to contain sequences that bind  $\alpha$ -scorpion toxins, which inactivate sodium channels, for use to evaluate ion channel activity and to screen for compounds for therapeutic applications. The modified sodium channel was then stably or transiently expressed in several mammalian host cells, including HEK293 variants and CHO cells, which were used in a high-throughput, plate-based screening assay.

International PCT publication WO/02068 (Korsgaard) describes stable cloning of a splice variant of a rat  $\alpha$ I sodium channel in HEK293 cells.

To date, there have been no reports of stable expression of a cloned human sodium type III channel in mammalian cells. The method described herein combines several procedures to facilitate the cloning and generation of stable cell

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lines containing such repetitive sequences, resulting in functional expression of such genes. In particular, the present invention describes the cloning and stable expression of a novel splice variant of human NaIII, designated hNaIII18.

The nucleotide sequence coding for hNaIII18, or an antigenic fragment, derivative or analog thereof, (including, *e.g.*, a chimeric protein) can be inserted into an appropriate expression vector, *i.e.*, a vector which contains the necessary elements for the transcription and translation of the inserted protein-coding sequence. Thus, a nucleic acid molecule having a nucleotide sequence encoding the hNaIII18 subunit protein of the invention can be operationally associated with a promoter in an expression vector of the invention. Either a cDNA or genomic sequence can be cloned and expressed under control of such regulatory sequences. Such vectors can be used to express functional, or functionally inactivated, hNaIII18 polypeptides.

The necessary transcriptional and translational signals can be provided on a recombinant expression vector, or they may be supplied from the native gene encoding hNaIII18 and/or its flanking regions.

Potential host-vector expression systems include but are not limited to mammalian cell systems transfected with expression plasmids or infected with virus (e.g., vaccinia virus, adenovirus, adeno-associated virus, herpes virus, etc.); insect cell systems infected with virus (e.g., baculovirus); microorganisms such as yeast containing yeast vectors; and bacteria transformed with bacteriophage, DNA, plasmid DNA, or cosmid DNA. The expression elements of vectors vary in their strengths and specificities. Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used.

Expression of the hNaIII18 protein may be controlled by any promoter/enhancer element known in the art, but these regulatory elements must be functional in the host selected for expression. Promoters which may be used to control hNaIII18 gene expression include, but are not limited to, cytomegalovirus (CMV) promoter (see, e.g., U.S. Patent Nos. 5,385,839 and 5,168,062), the SV40 early promoter region (Benoist and Chambon, Nature 1981; 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, et al., Cell, 1980; 22:787-797), the herpes thymidine kinase promoter (Wagner et al.,

Proc. Natl. Acad. Sci. U.S.A., 1981; 78:1441-1445), the regulatory sequences of the metallothionein gene (Brinster *et al.*, Nature, 1982; 296:39-42, prokaryotic expression vectors such as the β-lactamase promoter (Villa-Komaroff, *et al.*, Proc. Natl. Acad. Sci. U.S.A. 1978; 75:3727-3731), or the tac promoter (DeBoer, *et al.*, Proc. Natl. Acad. Sci. U.S.A. 1983; 80:21-25) (see also "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242:74-94), promoter elements from yeast or other fungi such as the Gal 4 promoter, the ADC (alcohol dehydrogenase) promoter, PGK (phosphoglycerol kinase) promoter, alkaline phosphatase promoter, and transcriptional control regions that exhibit tissue specificity, such as, *e.g.*, endothelial cell-specific promoters.

Solubilized forms of the protein can be obtained where necessary by solubilizing inclusion bodies or reconstituting membrane components, *e.g.*, by treatment with detergent, and if desired sonication or other mechanical processes, as described above. The solubilized protein can be isolated using various techniques, such as polyacrylamide gel electrophoresis (PAGE), isoelectric focusing, 2-dimensional gel electrophoresis, chromatography (*e.g.*, ion exchange, affinity, immunoaffinity, and sizing column chromatography), centrifugation, differential solubility, immunoprecipitation, by any other standard technique for the purification of proteins, or by a combination of such techniques.

Since  $\beta$ -subunits 1-3 are known to bind the  $\alpha$ -subunits of sodium channels, the present invention also contemplates co-expression of a  $\beta$ -subunit with NaIII18. While the role played by  $\beta$ -subunits in determining the pharmacological properties of voltage-gated sodium channels appears to be minor, at least for the commonly-studied binding sites,the  $\beta$ -subunits do appear to have effects on the biophysics (gating kinetics) of sodium channel function. Therefore, to the extent that biophysics and drug interactions are linked, the  $\beta$ -subunits may affect pharmacology of agents used to modulate sodium channel activity. Some known  $\beta$ -subunits that may be co-expressed with the NaIII18 subunit of the invention are described in Isom et al., Neuron 1994; 12:1183-94; International PCT publication WO 01/44293 to Plumpton et al.; International PCT publication WO 01/23570 to d'Andrea et al.; U.S. published patent application 2002/0045229 to Qin et al.; and under GenBank Accession Nos.

U87445, AF007783, AH005825, AF007783, AF04948, L10338 and L16242, among others

#### hNaIII18 Binding Partners

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The present invention further provides a method for identifying physiological binding partners of hNaIII18. One method for evaluating and identifying hNaIII18 binding partners is the yeast two-hybrid screen. Preferably, the yeast two-hybrid screen is performed using an cell library with yeast that are transformed with recombinant hNaIII18. Alternatively, hNaIII18 can be used as a capture or affinity purification reagent. In another alternative, labeled hNaIII18 can be used as a probe for binding, e.g., by immunoprecipitation or Western analysis. Several expected hNaIII18 binding partners are the sodium channel  $\beta$  subunits, as described in the section above.

Generally, binding interactions between hNaIII18 and any of its binding partners will be strongest under conditions approximating those found in the native cell, *i.e.*, physiological conditions of ionic strength, pH and temperature, and particularly those obtaining in the cell membrane. Perturbation of these conditions will tend to disrupt the stability of a binding interaction.

#### Antibodies to hNaIII18

Antibodies to hNaIII18 are useful, *inter alia*, for determining the presence of hNaIII18 in a cell and for cellular regulation (*i.e.*, inhibition) of hNaIII18 activity, as set forth below. According to the invention, a hNaIII18 polypeptide produced recombinantly or by chemical synthesis, and fragments or other derivatives or analogs thereof, including fusion proteins, may be used as immunogens to generate antibodies that recognize the hNaIII18 polypeptide. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments, and Fab expression libraries. Such an antibody binds specifically to hNaIII18, and may recognize either a mutant form of hNaIII18 or wild-type hNaIII18, or both. The antibodies of the present invention are specific for hNaIII18 and either do not recognize, or bind with lower affinity to, orthologs of hNaIII18. In one embodiment,

specific binding of such antibodies to hNaIII18 polypeptides provides the ability to detect the presence of the hNaIII18 polypeptide in a sample. In another embodiment, specific binding of such antibodies to hNaIII18 polypeptides provides the ability to preferentially inhibit the activity of hNaIII18, or an ion channel comprising hNaIII18.

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Various procedures known in the art may be used for the production of antibodies against hNaIII18 polypeptides. These include but are not limited to the hybridoma technique originally developed by Kohler and Milstein (Nature 1975; 256:495-497), as well as the trioma technique, the human B-cell hybridoma technique (Kozbor *et al.*, Immunology Today 1983, 4:72; Cote *et al.*, Proc. Natl. Acad. Sci. 1983, 80:2026-2030), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole *et al.*, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., 1985, pp. 77-96).

### hNaIII18 Agonists and Antagonists

The present invention also contemplates the identification of compounds that modulate hNaIII18 sodium channel activation and activity. Such compounds are useful, *e.g.*, for inhibiting (*i.e.*, antagonizing) or increasing (*i.e.*, agonizing) biological activities that are associated with sodium channel activation and/or as therapeutic agents for treating disorders associated with excessive sodium channel activation.

Compounds that modulate hNaIII18 activity or an activity associated therewith may be readily identified using screening methods of the present invention. In one embodiment, compounds identified by the screening methods of this invention bind to a hNaIII18-subunit containing ion channel. Compounds identified by the present method may antagonize or agonize hNaIII18 subunit-containing channel activity, as well as a related downstream biological effect (*e.g.*, the ability of DRG to transmit nociceptive signals from the PNS to the CNS) that are associated with excessive sodium channel current and activity.

In vivo or cell culture assays may be used to determine whether a test compound functions as an antagonist to inhibit hNaIII18 activity in cells. For instance, cell culture assays may be used to measure a test compound's ability to modulate an activity, such as induction, strength or duration of sodium channel

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current associated with hNaIII18 subunit-containing sodium channel activity. Such assays generally comprise contacting a cell that expresses a hNaIII18 subunit containing sodium channel with a test compound. The cell should preferably be contacted with the test compound before or during exposure to an agent or stimulus that otherwise would serve to depolarize the cell membrane and thus activate (i.e., open) the sodium channel: e.g. a high potassium chloride saline solution, or an extracellular field-stimulating electrode. The cell can then be examined to determine whether a response otherwise associated with sodium channel activation has been inhibited. In a non-limiting embodiment, the response of the cell treated with the test compound is compared to that of a control cell that has not been treated with the test compound. Cell assays include those utilizing conventional, electrode-based, electrophysiological techniques, as well as the new generation high-throughput, planar electrode (orifice) -based, electrophysiological technologies, among others. Other assays include monitoring changes in membrane potential with appropriate fluorescent, or luminescent, dyes, measuring ion flux through the sodium channel with a radiolabeled tracer, or assaying downstream consequences of sodium channel activation, such as calcium mobilization or effects on gene expression, using an appropriate reporter system.

Positive modulation (*i.e.*, agonism) of hNaIII18 subunit-containing channels may be desirable under certain circumstances, and screening for such agonists can be conducted according to the methods of the invention.

#### Screening

According to the present invention, nucleotide sequences encoding hNaIII18 are useful targets to identify drugs that are effective in preventing or alleviating pain, or drugs that can be used as anti-epileptics/anticonvulsants, anesthetic antiarrythmics, and in the treatment of bipolar disorder (see section entitled Therapeutics, below), any of which may be associated with the function of the sodium channel. Examples of such drugs include without limitation: (i) isolated nucleic acids capable of altering expression of hNaIII18 (e.g., antisense or ribozyme molecules); (ii) small organic molecules that bind to and modulate the function of a hNaIII18 subunit or a hNaIII18 subunit-containing ion channel; and (iii) peptides or

peptide analogs that bind to and modulate the function of a hNaIII18 subunit or a hNaIII18 subunit-containing ion channel. In addition, the nucleotide sequences encoding hNaIII18 are useful for studying the role of the channels both in pain perception and in physiological and pathological brain functions.

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Any screening technique known in the art can be used to screen for agonists or antagonists. The present invention contemplates screens for small molecules and mimics, as well as screens for natural products that bind to and agonize or antagonize hNaIII18-containing ion channels. For example, natural product libraries can be screened using assays of the invention for molecules that agonize or antagonize hNaIII18-containing ion channel activity.

Knowledge of the primary sequence of hNaIII18, and the similarity of that sequence with proteins of known function, can provide an initial lead to inhibitors or antagonists. Identification and screening of modulators is further facilitated by determining structural features of the protein, *e.g.*, using X-ray crystallography, neutron diffraction, nuclear magnetic resonance spectrometry, and other techniques for structure determination. These techniques provide for the rational design or identification of agonists and antagonists.

Another approach uses recombinant bacteriophage to produce large libraries. Using the "phage method" (Scott and Smith, Science 1990, 249:386-390; Cwirla, et al., Proc. Natl. Acad. Sci. USA 1990, 87:6378-6382; Devlin et al., Science 1990, 49:404-406), very large libraries can be constructed (106-108 chemical entities). A second approach uses primarily chemical methods, of which the Geysen method (Geysen et al., Molecular Immunology 1986, 23:709-715; Geysen et al. J. Immunologic Methods 1987, 102:259-274); and the method of Fodor et al. (Science 1991, 251:767-773) are examples. Furka et al. (14th International Congress of Biochemistry 1988, Volume #5, Abstract FR:013; Furka, Int. J. Peptide Protein Res. 1991, 37:487-493), Houghton (U.S. Patent No. 4,631,211) and Rutter et al. (U.S. Patent No. 5,010,175) generally describe methods to produce a mixture of peptides that can be tested as agonists or antagonists.

In another aspect, synthetic libraries, such as those described in Needels et al., Proc. Natl. Acad. Sci. USA 1993, 90:10700-4; Ohlmeyer et al., Proc. Natl. Acad. Sci. USA 1993, 90:10922-10926; Lam et al., PCT Publication No. WO

92/00252; and Kocis et al., PCT Publication No. WO 9428028, and the like, can be adapted to screen for compounds according to the present invention.

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Test compounds can be screened from large libraries of synthetic or natural compounds. Numerous means are currently used for random and directed synthesis of saccharide, peptide, and nucleic acid based compounds. Synthetic compound libraries are commercially available from a variety of sources, including Maybridge Chemical Co. (Trevillet, Cornwall, UK), Comgenex (Princeton, NJ), Brandon Associates (Merrimack, NH), and Microsource (New Milford, CT). A rare chemical library is available from Aldrich (Milwaukee, WI). Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available from a variety of sources including, *e.g.*, Pan Laboratories (Bothell, WA) and MycoSearch (NC), or are readily producible de novo. Additionally, natural and synthetically produced libraries and compounds are readily modified through conventional chemical, physical, and biochemical means (see, *e.g.*, Blondelle et al., TIBTech 1996, 14:60).

#### In Vitro Screening Methods and Activity Assays

#### **Cell-based screening**

Intact cells expressing a hNaIII18 subunit-containing ion channel can be used in screening methods to identify candidate compounds useful in modulating the activity of sodium channels containing hNaIII18. In one embodiment, a cell line is established that stably expresses or overexpresses the hNaIII18 subunit protein, either alone or in combination with one or more other sodium channel  $\beta$  subunits, to form a functional sodium channel. Alternatively, cells (including without limitation mammalian, invertebrate, yeast, or bacterial cells) are transiently programmed to express a hNaIII18 subunit protein by introduction of the appropriate DNA or mRNA. Identification of candidate compounds can be achieved using any suitable assay, including without limitation: (i) assays that measure binding of test compounds to hNaIII18 (alone or in combination with sodium channel  $\beta$  subunits described *supra*): (ii) assays that measure the ability of a test compound to modulate (*i.e.*, agonize or antagonize) a measurable activity or function of hNaIII18 or a hNaIII18 subunit-containing ion channel; and (iii) assays that measure the ability of a compound to

enhance or inhibit the transcriptional activity of sequences derived from the promoter (*i.e.*, regulatory) regions of the hNaIII18 gene.

Any cell assay system that allows for assessment of functional activity of a hNaIII18 subunit-containing sodium channel is encompassed by the present invention. In a specific embodiment, described *infra*, the assay can be used to identify compounds that selectively modulate the hNaIII18 subunit protein, which can be determined by assessing the effects on NaIII18 subunit-expressing cells contacted with a test compound. The assay system can thus be used to identify compounds that selectively produce a functional effect through hNaIII18 sodium channels.

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Compounds that decrease activity of the sodium channel in response to activation may be useful as novel therapeutics in the amelioration of neuropathic pain mediated by DRG neurons, or as anti-epileptics/convulsants, anesthetics, antiarrythmics, or in the treatment of bipolar disorder.

Compounds that increase activity of sodium channels may be useful as cognitive enhancers, or in disorders such schizophrenia. In these instances, a subtype-selective agent would be preferable to offset the potential for proconvulsant effects and to increase cardiac contractility in individuals suffering from heart failure.

Alternatively, the change in membrane potential induced by sodium ions of the voltage-gated channel-containing cells may be monitored using fluorescence methods. When using fluorescence methods, the voltage-gated channel containing cells may be incubated with a membrane potential indicating agent that allows for a determination of changes in the membrane potential of the cells caused by the influx of sodium ions. Such membrane potential indicating agents include fluorescent indicators, such as those provided in a Molecular Devices Membrane Potential Kits for the FLIPR/Flexstation, DIBAC4(3), DiOC6(6) DiOC5(3), DiOC2(3) and fluorescence resonance energy transfer (FRET) based dyes such as JC1, and JC9, among others.

Another method that allows for assessment of functional activity of hNaIII18-containing sodium channels involves monitoring the change in membrane potential induced by sodium ions on the channel-containing cells by fluorescent methods, *e.g.*, using a FLIPR assay (Fluorescence Image Plate Reader; available from Molecular Devices)(Rose et al. Pflugers Arch. 1999 Dec;439(1-2):201-7). Another

method involves radioactive flux assays that measure the ability of radioactive tracer ions such as [<sup>22</sup>Na] and [<sup>14</sup>C] guanidinium to pass into the cell upon channel activation (Barann M. et al. Naunyn Schmiedebergs Arch Pharmacol. 1999; 360(3):234-41). After the channel is activated, concentrations of these tracer ions increase inside the cell. Free extra-cellular tracer is washed away, cells are lysed, and radioactivity in the lysates is counted using standard scintillation counters or other radioactivity analysis instruments.

Yet another method involves measuring cell viability upon veratridine-mediated stabilization of sodium channels in their open conformation (Okuyama K. et al., Eur J Pharmacol. 2000; 398(2):209-16). Cells undergo toxic sodium overload followed by cell death. Compounds that prevent cell death, or cellular toxicity, can be assayed with standard cytoxicity kits and with standard cell viability dyes such as alamar blue.

Cell-Free Screening

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In another embodiment, an assay is a cell-free assay comprising contacting a hNaIII18 polypeptide or biologically active portion thereof with a test compound and determining the ability of the test compound to modulate (e.g., stimulate or inhibit) the activity of the hNaIII18 polypeptide or biologically active portion thereof.

In yet another embodiment, the cell-free assay comprises (i) contacting the hNaIII18 polypeptide of the invention or biologically active portion thereof with a known compound or polypeptide which binds the hNaIII18 polypeptide to form an assay complex; (ii) contacting the assay complex with a test compound; (iii) determining the ability of the test compound to interact with the hNaIII18 polypeptide by determining the ability of the test compound to modulate the effect of the known compound on the activity of the sodium channel.

More specifically, a cell-free method can involve monitoring the specific binding of a radiolabeled sodium channel selective neurotoxin, such as [<sup>3</sup>H]tetrodotoxin or [<sup>3</sup>H]batrachotoxin, or a high affinity small-molecule ligand, to a membrane preparation from cells or tissues engineered to express hNaIII18-containing sodium channels (Garritsen A. et al. Eur J Pharmacol. 1988; 145(3):261-6;

MacKinnon AC. et al. J Pharmacol. 1995; 115(6):1103-9; Bambrick L. et al., J Pharmacol Toxicol Methods. 1994; 32(3):129-38). Following techniques that are well know in the art, total binding to membranes can be measured upon incubation with the radioligand until the biomolecular reaction reaches equilibrium. Nonspecific binding is defined in the presence of an unlabelled competitor ligand. Specific binding is the subtraction of total minus nonspecific binding. Compounds that modulate specific binding can thereby be identified.

In another embodiment, modulators of expression of the hNaIII18 polypeptide of the invention are identified in a method in which a cell is contacted with a candidate compound and the expression of the mRNA or protein corresponding to hNaIII18 in the cell is determined. The level of expression of the hNaIII18 mRNA or protein in the presence of the candidate compound is compared to the level of expression of the hNaIII18 mRNA or protein in the absence of the candidate compound. The candidate compound can thereby be identified as a modulator of expression of the hNaIII18 polypeptide of the invention based on this comparison. For example, when expression of the hNaIII18 mRNA or protein is increased in the presence of the candidate compound compared to in the absence of the candidate compound, then the candidate compound is identified as a stimulator of hNaIII18 mRNA or protein expression. Alternatively, when expression of the hNaIII18 mRNA or protein is specifically reduced in the presence of the candidate compound compared to in the absence of the candidate compound, then the candidate compound is identified as an inhibitor of hNaIII18 mRNA or protein expression. In view of this disclosure, the level of the hNaIII18 mRNA or protein expression in cells can be determined by methods known in the art.

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#### High-Throughput Screen

Drug candidates according to the invention can be identified by screening in high-throughput assays, including without limitation cell-based or cell-free assays. It will be appreciated by those skilled in the art that different types of assays can be used to detect different types of drug candidates. Several methods of automated assays have been developed in recent years so as to permit screening of tens of thousands of compounds in a short period of time. Such high-throughput

screening methods are particularly preferred. The use of high-throughput screening assays to test for agents is greatly facilitated by the availability of the large amounts of purified hNaIII18 polypeptides provided by the invention.

#### Therapeutic Uses

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It is desirable to modulate the function of sodium channels in a number of clinical and therapeutic environments. Sodium channels are implicated in conditions including chronic and neuropathic pain, cardiac arrhythmias (Duch et al., Toxicol Lett 1998; 100-101:255-63), neuronal disorders associated with deficient oxygen supply or mitochondrial dysfunction (Urenjak et al., Amino Acids 1998;14(1-3):151-8), and epilepsy (Ragsdale et al., Brain Res Rev 1998;26(1):16-28). In addition, inhibition of sodium channels is an effect of local anesthetics (Li et al., Mol Pharmacol 1999; 55(1):134-41).

According to the present invention, inhibition of hNaIII18 subunit-containing sodium channel activity may be used as a treatment option in patients with a pain disorder, such as but not limited to a neuropathic pain-related disease such as, *e.g.*, pain from peripheral nerve trauma, herpes virus infection, diabetes mellitus, causalgia, plexus avulsion, neuroma, limb amputation, and vasculitis. Neuropathic pain is also caused by nerve damage from chronic alcoholism, human immunodeficiency virus infection, hypothyroidism, uremia, or vitamin deficiencies. The neuronal hyperexcitability and corresponding molecular changes in neuropathic pain have many features in common with the cellular changes in certain forms of epilepsy. This has led to the use of anticonvulsant drugs for the treatment of neuropathic pain (Jensen, Eur J Pain 2002;6 Suppl A:61-8). Local anesthetics such as lidocaine and mexiletine have also be shown to inhibit TTX-S sodium channel activity in hyperexcitable neurons in rat (Novartis Found Symp 2002;241:189-201; discussion 202-5, 226-32).

Inhibition of the sodium channel of the present invention may also be used as a treatment option in patients with chronic pain. In chronic pain, the pain can be mediated by multiple mechanisms. This type of pain generally arises from injury to the peripheral or central nervous tissue. The chronic pain-type syndromes include pain associated with spinal cord injury, multiple sclerosis, post-herpetic neuralgia,

trigeminal neuralgia, phantom pain, causalgia, and reflex sympathetic dystrophy and lower back pain.

Inhibition of the sodium channel of the present invention may also be used as a treatment option in patients with nociceptive pain.

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### Inhibition of Protein Synthesis or Sodium Channel Activity

Gene transcription and protein translation may be inhibited by administration of exogenous compounds. Exogenous compounds may interact with extracellular and/or intracellular messenger systems to regulate protein synthesis. In this embodiment, exogenous compounds that inhibit hNaIII18 protein synthesis may be used in the prevention and/or treatment for pain resulting from persistent channel activity.

Accordingly, in an exemplary embodiment, the modulatory method of the invention involves contacting a cell, tissue or subject with an agent that modulates one or more of the activities of hNaIII18 protein activity associated with the cell. An agent that modulates hNaIII18 protein activity can be an agent as described herein, such as a nucleic acid or a protein, an hNaIII18-specific antibody, an hNaIII18 agonist or antagonist, a peptidomimetic of an hNaIII8 agonist or antagonist, or other small molecule. In one embodiment, the agent stimulates one or more hNaIII18 activities. In another embodiment the agent inhibits one or more hNaIII18 activities. Examples of such inhibitory agents include antisense hNaIII18 nucleic acid molecules, antihNaIII18 antibodies, and hNaIII18 inhibitors. These modulatory methods can be performed in vitro (e.g., by culturing the cell with the agent) or, alternatively, in vivo (e.g., by administering the agent to a subject). As such, the present invention provides methods of treating an individual afflicted with a disease or disorder characterized by aberrant or unwanted expression or activity of a hNaIII18 protein or nucleic acid molecule. In one embodiment, the method involves administering an agent (e.g., an agent identified by a screening assay described herein), or combination of agents that downregulates hNaIII18 expression or activity or the activity of a hNaIII18 subunitcontaining ion channel.

In yet another embodiment, the agent enhances one or more hNaIII18 activities, such as by administering a hNaIII18 protein or nucleic acid molecule as therapy to compensate for reduced or aberrant hNaIII18 expression or activity.

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The present invention further provides antisense nucleic acids, which may be used to inhibit expression of hNaIII18 nucleotide sequences of the invention. This antisense technology has been described as inhibiting the peripheral tetrodotoxin (TTX)-resistant sodium channel, NaV1.8, found in sensory neurons, when administered intrathecally (Lai et al., Pain 2002; 95 (1-2):143-52). According to this method, the antisense nucleic acid, upon hybridizing under cytoplasmic conditions with complementary bases in an RNA or DNA molecule, inhibits the RNA or DNA. Additionally, hybridization of the antisense nucleic acid to the DNA or RNA may inhibit transcription of the DNA into RNA and/or translation of the RNA into the protein. If the RNA is a messenger RNA transcript, the antisense\_nucleic acid is a counter-transcript or mRNA-interfering complementary nucleic acid. Antisense nucleic acid molecules can be encoded by a recombinant gene for expression in a cell (see, e.g., U.S. Patent No. 5,814,500; U.S. Patent No. 5,811,234) or can be prepared synthetically (e.g., U.S. Patent No. 5,780,607).

Alternatively, antibody molecules or antigen-binding antibody fragments can be administered either directly or by expressing nucleotide sequences encoding antibodies or binding fragments thereof within the target cell population by utilizing, for example, techniques such as those described in Marasco *et al.* (Proc. Natl. Acad Sci. USA, 1993, 90:7889-7893).

#### Formulations and Administration

The drug candidate or agent that modulates hNaIII18 activity is advantageously formulated in a pharmaceutical composition by admixing the drug candidate or agent with a pharmaceutically acceptable carrier. This agent may then be designated as the active ingredient, or therapeutic agent for use, for example, against chronic, neuropathic pain, or nociceptive pain

The form, amount and route of administration of the therapeutic compound envisioned for use depends on the type and severity of the disease or condition to be treated, as well as the patient's state of health, gender, weight, age,

etc., and can be determined by an attending medical practitioner in view, e.g., of the results of published clinical trials. The concentration or amount of the active ingredient depends on the desired dosage and administration regimen, as discussed below. Suitable dose ranges may include from about 1 mg/kg to about 100 mg/kg of body weight per day.

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The pharmaceutical compositions may also include other biologically active substances in combination with the NaIII18 modulatory agent. Such substances include but are not limited to opioids such as morphine, codeine, fentynyl, oxycodone, hydrocodone, and buprenorphine; and non-steroidal anti-inflammatory drugs (NSAID's) such as but not limited to ibuprofen and COX-2 inhibitors, among others

The phrase "pharmaceutically acceptable" refers to molecular entities and compositions that are physiologically tolerable and do not typically produce an allergic or similar untoward reaction, such as gastric upset, dizziness and the like, when administered to a human. Preferably, as used herein, the term "pharmaceutically acceptable" means that the carrier has been approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, and more particularly in humans. The term "carrier" refers to a diluent, adjuvant, excipient, or vehicle with which the active ingredient is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. Water or aqueous solution saline solutions and aqueous dextrose and glycerol solutions are preferably employed as carriers, particularly for injectable solutions. Suitable pharmaceutical carriers are described in "Remington's Pharmaceutical Sciences" by E.W. Martin.

According to the invention, the pharmaceutical composition of the invention can be introduced parenterally, transmucosally, *e.g.*, orally (per os), nasally, rectally, or transdermally. Parental routes include intravenous, intra-arteriole, intramuscular, intradermal, subcutaneous, intraperitoneal, intraventricular, and intracranial administration. The pharmaceutical composition may alternatively be

adapted for topical or transdermal application, such in a salve, cream, lotion, spray or transdermal patch system.

The pharmaceutical compositions may be added to a retained physiological fluid such as blood or synovial fluid. For CNS (Central Nervous System) administration, a variety of techniques are available for promoting transfer of the therapeutic across the blood brain barrier including disruption by surgery or injection, co-administration of drugs that transiently open adhesion contact between CNS vasculature endothelial cells, and co-administration of substances that facilitate translocation through such cells.

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In another embodiment, the active ingredient can be delivered in a vesicle, in particular a liposome (see Langer, Science 1990; 249:1527-1533; Treat et al., in Liposomes in the Therapy of Infectious Disease and Cancer, Lopez-Berestein and Fidler (eds.), Liss: New York 1989 pp. 353-365; Lopez-Berestein, ibid., pp. 317-327; see generally ibid.).

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In yet another embodiment, the therapeutic substance can be delivered in a controlled release formulation. For example, an active ingredient may be administered using intravenous infusion with a continuous pump, in a polymer matrix such as poly-lactic/glutamic acid (PLGA), a pellet containing a mixture of cholesterol and the active ingredient (SilasticRTM; Dow Corning, Midland, MI; see U.S. Patent No. 5,554,601) implanted subcutaneously, an implantable osmotic pump, a transdermal patch, liposomes, or other modes of administration.

Compounds identified in the screening methods described herein (*i.e.*, modulators of sodium channel activity), may be provided to the patient in formulations that are known in the art and may include any pharmaceutically acceptable additives, such as excipients, lubricants, diluents, flavorants, colorants, and disintegrants. The formulations may be produced in useful dosage units such as tablet, caplet, capsule, liquid, or injection. In a further embodiment, these compounds are also administered in conjunction with other therapeutic agents such as the local anesthetics and anti-epileptic or anti-convulsants discussed *supra*.

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The form and amount of therapeutic compound envisioned for use depends on the type of disease and the severity of the desired effect, patient state, etc., and can be determined by one skilled in the art.

#### **EXAMPLES**

The present invention is also described by means of an example, presented below. The use of such an example is illustrative only and in no way limits the scope and meaning of the invention or of any exemplified term. Likewise, the invention is not limited to any particular preferred embodiments described herein. Indeed, many modifications and variations of the invention will be apparent to those skilled in the art upon reading this specification and can be made without departing from its spirit and scope. The invention is therefore encompassed by the appended claims along with the full scope of equivalents to which the claims are entitled.

# EXAMPLE 1: CLONING AND EXPRESSION OF HUMAN NaIII18 Methods

Reverse transcription and amplification of hNaIII18 cDNA. Reverse transcription was carried out using ThermoScript Reverse Transcriptase (Life Technologies, Rockville, MD), at an annealing temperature of 55 °C to maximize the likelihood of obtaining a full-length mRNA, according to manufacturer's instructions.

The following primers were designed to amplify the resulting full-length hNaIII18 cDNA:

forward	5' - ATAAGAATGCGGCCGCTGAAAAGATGGCACAGGCAC-3'
primer (SEQ	
ID NO: 7)	-
reverse	5' - ATAGTTTAGCGGCCGCCTTGAAGTCCAGTTGACACA -3'
primer (SEQ	- 1111101111111111111111111111111111111
ID NO: 8)	

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Primers were designed from the human NaIII (SCN3A) mRNA sequence previously identified (GenBank Accession # AJ251507).

Full-length cDNA (6000 base-pairs) was amplified using the Expand Long Template PCR (Boehringer Mannheim, Indianapolis, IA) according to the manufacturer's instructions. This enzyme is a mixture of thermostable Taq and Pwo

DNA polymerases. The number of cycles used for amplification was decreased to 28 cycles instead of the traditional 30-35 as an added precaution to minimize the occurrence of mutations during PCR.

Purification and cloning of PCR products into expression vectors.

PCR products resulting from the above-described reaction were visualized after electrophoresis on an agarose gel containing Crystal Violet. DNA was purified from the gel using methods well known in the art. DNA was stored in Tris-EDTA buffer, pH 7.4.

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The PCR-amplified cDNA was cloned into a low-copy number expression vector, pLCTM1 (kindly provided by Al Goldin, UCI) according to standard procedures. This vector is under the control of the origin of replication (ORI) from plasmid pACYC184, which has a limited number of replication cycles, resulting in a decreased error rate during DNA replication.

Further, the plasmid contains a tetracycline-resistance gene instead of an ampicillin-resistance gene for selection. Tetracycline is less likely to induce mutations than ampicillin during selection. The plasmid also contains a neomycin resistant gene (NeoR) for selection of stable cell lines using the neomycin analog G418.

Once cloned, the vectors were transformed into maximum efficiency STBL2 competent *E. coli* bacteria (Life Technologies, Rockville, MD), provided in the kit according to manufacturer's instructions. These cells optimize the cloning of unstable inserts. Bacteria expressing hNaIII18 were grown at 30-33°C, and maintained in exponential (log) growth phase for the duration of culture.

Small tetracycline-resistant colonies were selected and grown-up for small-scale DNA preparations and large-scale preparations. The concentration of tetracycline was kept low (15  $\mu$ g/ml) to further minimize adverse growth conditions. The cDNA was extracted using the Wizard Plus SV Minipreps DNA Purification System Kit (Promega, Madison, WI) according to the manufacturer's instructions, or Qiagen Midipreps according to manufacturer's instructions (Qiagen, Valencia, CA). cDNA was then analyzed by restriction digest, and partial sequencing. Full sequencing was performed by MWG (North Carolina). Partial sequencing was done with standard DTCS sequencing method using a commercial Beckman Coulter kit.

Clones, human embryonic kidney cells (HEK293) were transiently transfected with clones that were identified as having the correct insert, and surveyed by an electrophysiological assay (Fugene transfection reagent, according to manufacturer's recommendation). One clone, pLCTM1huNaIII-18, was determined to be functional as it gave large TTX-S currents with the expected activation and inactivation kinetics typical of NaIII channel. For example, typical activation is measured within fractions of ms at Vm=0mV (corresponding Imax). Inactivation is measured as the time constant between 1-3 ms at Vm=0mV (increasing to 20 ms at -50 mV to 0.5 ms at +40mV). Recovery from inactivation is a time constant of about 10ms at Vm=-100mV and 60 ms at -80mV (see e.g., Cummins et al., J Neurosci 2001; 21:52-5961).

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This clone was fully sequenced for confirmation. In addition, several non-functional clones were partially sequenced.

Clone pLLCTM1huNaIII-18 was used to generate a stable cell line in HEK293 cells. Fugene-mediated transfection of HEK cells was performed in 35 mm dish followed by G418 selection (300 and 500  $\mu$ g/ml), colony isolation, line expansion. G418-resistant cells were then analyzed with immunocytochemistry, RT-PCR and electrophysiology according to standard techniques.

Electrophysiology. Stably transfected cells were grown on poly DL-lysine-coated glass coverslips at ~2,000 cells/slip, or Petri dishes at ~10,000 cells/dish and were then placed into the electrophysiology recording chamber and infused with an extracellular solution (140 mM NaCl, 4.7 mM KCl, 1.2 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 11 mM glucose and 5 mM HEPES, pH 7.4) at a rate of 2 ml/min. Electrodes were prepared by pulling Patch pipettes (borosilicate glass) using a Sutter P-97 electrode puller, and were filled with a solution containing 110 mM CsCl, 10 mM NaCl, 5 mM MgCl<sub>2</sub>, 11 mM EGTA, 10 mM HEPES, 2 mM ATP and 1 mM GTP, pH 7.25, osmolarity 275-290 mOsm. When filled with this solution, the electrodes had resistances of about 1-4 MS. Currents were recorded using a whole-cell voltage clamp techniques as described in Hamill et al. (Pflugers Arch. 1981; 391; 85-100), at room temperature (21-23 °C). Briefly, currents were recorded using an Axopatch 200A amplifier (Axon Instruments, Foster City, CA) and were leak-subtracted (P/4),

low-pass filtered (3 kHz, 8-pole Bessel), digitized (20-50- $\mu$ s intervals), and stored using Digidata 1200 B interface and Pclamp6/Clampex software (Axon Instruments, Foster City, CA). Residual series access resistance was largely (75-80%) canceled using built-in amplifier circuitry. The junction potential calculated using JPCalcW software (Cell MicroControl, Virginia Beach, VA) was small (<7 mV); so, no correction of the holding voltage was made.

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To take I-V curves, cells were held at a holding voltage,  $V_h = -90 \text{mV}$ . A series of 16 depolarizing pulses (10ms in duration) incrementing in 10 mV steps were applied at a frequency of 0.5 Hz. The peak values of currents were plotted against corresponding voltage steps to get the I-V curve. From this plot  $V_{max}$ , *i.e.*, the voltage causing the maximal  $Na^+$  current, as well as rising times to peak and time constant for inactivation at different voltages were determined. To get steady-state inactivation curves, cells were held at a holding voltage,  $V_h = -120 \text{mV}$  to remove residual inactivation. A series of 30 depolarizing conditioning pre-pulses (each 100ms in duration) incrementing in 5 mV steps immediately followed by a 5 ms testing pulse,  $V_t$ , to  $V_{max}$  were applied at a frequency of 0.5 Hz. The peak currents in response to  $V_t$  were plotted against the size of corresponding conditioning pre-pulses,  $V_c$ , to get steady-state inactivation curve. The Boltzman fit to this curve, *i.e.*,  $\{1/[1+\exp((V+V/2)/k)]\}$ , returned the values of  $V/_2$  (the half-inactivation voltage) and k (the slope of the curve).

To measure recovery from inactivation, cells were held at a holding voltage

 $V_h$ = -120mV to remove residual steady-state inactivation. The depolarizing conditioning pre-pulse (100 ms in duration) was applied to  $V_c$  to cause complete inactivation of the channels (usually  $V_c$ =-10 mV). The conditioning pre-pulse was immediately followed by hyperpolarizing gap back to -120mV of a variable duration. The gap duration was incremented in subsequent cycles in varying steps (2 ms -100 ms) depending on the speed of recovery. The gap was immediately followed by the testing pulse  $V_t$  (10 ms in length) to assess the fraction of  $Na^+$  channels available for activation. The cycle was repeated every 5 seconds while the gap duration was incremented. The peak currents to  $V_t$  were plotted against the corresponding gap

duration to get the kinetics of recovery. The mono- or double- exponential fit to the data returned the time constant,  $\tau_{repr.}$ , of repriming from inactivation.

#### Results

Identification of a splice-variant for human NaIII (SCN3). Clone

pLCM1huNaIII-18 is a novel splice variant and contains an additional 147 nucleotides corresponding to 49 amino acids in the cytoplasmic loop between domain 1S6 and IIS1 (see SEQ ID NO: 1 and SEQ ID NO: 2). Partial sequencing of several other clones that were not determined to have functional activity revealed sequences that either matched the published sequence (GenBank Accession #AJ251507) or

what had been described for the rat NaIII clone (Schaller et al., *J Neurosci* 1992; 12(4):1370-81), resulting in a protein with an additional 3 (rNaIIIa) or 22 (rNaIIIb)

contained an extra 9 or 96 nucleotides. The shorter splicing patterns correspond to

amino acids, but had not been described for the human NaIII before.

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Subsequent to the completion of the cloning of hNaIII18, it was discovered that a clone having the same 147 nucleotide insert was deposited in GenBank on February 1, 2001 (GenBank Accession # AF225986-SEQ ID NO: 5). See cDNA alignment in Figure 8. However, that encoded amino acid sequence differs from the sequence disclosed herein by 12 amino acids (between two clones), at amino acid residues 208, 475, 495, 508, 604,1163, 1576, 1614, 1741, 1743, 1862 and 1966, respectively (SEQ ID NO: 2 vs. SEQ ID NO: 6). See amino acid alignent of Figure 9.

Stable transfection of the pLCM1huNaIII-18 resulted in the generation of two cell lines that expressed the expected ~220 kDa hNaIII18 protein and exhibited functional sodium channels, designated 293/huNaIII18-300-20 and 293/huNaIII18-500-35, with appropriate TTX-S currents. 293/huNaIII18-300-20 had an activation threshold voltage of -40 mV (Figure 9A), a steady state V ½ inactivation voltage of -58 mV (Figure 9B), a recovery time after inactivation of 2.5 ms (fast component) AND 113 ms (slow component-(Figure 9C), and inactivation kinetics of 0.8 ms (Figure 9D).

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The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and the accompanying figures. Such modifications are intended to fall within the scope of the appended claims.

Patents, patent applications, publications, procedures, and the like are cited throughout this application, the disclosures of which are incorporated herein by reference in their entireties.

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#### WHAT IS CLAIMED IS:

- 1. An isolated nucleic acid comprising a nucleotide sequence encoding a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO: 2).
- 2. The isolated nucleic acid of claim 1, comprising the nucleotide sequence of Figure 1 (SEQ ID NO: 1).
- 3. A recombinant vector comprising a nucleotide sequence encoding a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO:2).
  - 4. A host cell comprising the recombinant vector of claim 3.
  - 5. A host cell genetically engineered to comprise the nucleic acid of claim 1.
  - 6. The host cell of claim 5 which is eukaryotic.
- 7. A eukaryotic host cell genetically engineered to express, or overexpress, a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO: 2).
- 8. A method for expressing a polypeptide in a cell cultured *in vitro* comprising culturing the cell of claim 4, 5, 6 or 7 under conditions conducive to the expression of the polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).
- 9. An isolated polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).

10. A host cell genetically engineered to co-express a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2) and a  $\beta$ -subunit of a sodium channel selected from the group consisting of  $\beta$ 1,  $\beta$ 2, and  $\beta$ 3.

- 11. An antibody or antigen-binding fragment that specifically binds to a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO: 2).
  - 12. The antibody of claim 11, which is a monoclonal antibody.
- 13. A method for detecting expression in a sample of a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises detecting specific binding of the antibody or antigen-binding fragment of claim 11 to a polypeptide in the sample.
- 14. A method for identifying a test compound that binds to a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises:
- (i) contacting a host cell that expresses a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2) with a test compound; and
- (ii) determining whether the test compound binds to the host cell but not to a control cell that does not express a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).
- 15. An assay method for identifying a test compound that modulates the activity of a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises:
- (i) providing a host cell that expresses a functional sodium channel comprising at least one polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2),
- (ii) contacting the host cell with a test compound under conditions that would activate sodium channel activity of said functional sodium channel in the absence of

the test compound; and

(iii) determining whether the host cell contacted with the test compound exhibits a modulation in activity of the functional sodium channel.

- 16. The assay method of claim 15, wherein the host cell has been genetically engineered to express or overexpress the functional sodium channel.
- 17. The assay method of claim 15, wherein the host cell has been genetically engineered by the introduction into the cell of a nucleic acid molecule having a nucleotide sequence encoding a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).
- 18. The assay method of claim 15, wherein the host cell has been genetically engineered to upregulate the expression of a nucleic acid encoding a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2),
- 19. The assay method of claim 18, wherein the upregulated nucleic acid is endogenous to the host cell.
- 20. The assay method of claim 15, wherein the modulation of the functional sodium channel activity is antagonism of that activity.
- 21. The assay method of claim 15, wherein the modulation of the functional sodium channel activity is agonism of that activity.

### FIGURE 1: NaIII18 cDNA (SEQ ID NO: 1)

tgaaaagatgcacaggcactgttggtacccccaggacctgaaagcttccgcctttttactaga gaatctcttgctgctatcgaaaaacgtgctgcagaagagaaagccaagaagcccaaaaaggaac aagataatgatgatgagaacaaaccaaagccaaatagtgacttggaagctggaaagaaccttcc atttatttatggagacattcctccagagatggtgtcagagcccctggaggacctggatccctac tatatcaataagaaaacttttatagtaatgaataaaggaaaggcaattttccqattcaqtqcca cctctgccttgtatattttaactccactaaaccctgttaggaaaattgctatcaagattttggt acattetttatteageatgettateatgtgeaetattttgaeeaaetgtgtatttatgaeettg agcaaccctcctgactggacaaagaatgtagagtacacattcactqgaatctatacctttqaqt cacttataaaaatcttggcaagaggttttgcttagaagattttacgtttcttcgtgatccatg gaactggctggatttcagtgtcattgtgatggcgtatgtaacagaatttgtaagcctaggcaat gtttcagcccttcgaactttcagaqtcttqagaqctctqaaaactatttctqtaattccaqqtt taaagaccattgtgggggccctgatccagtcggtaaagaagctttctgatqtqatqatcctqac tgtgttctgtctgagcgtgtttgctctcattgggctgcagctgttcatgggcaatctgaggaat aaatgtttgcagtggcccccaagcgattctgcttttgaaaccaacaccacttcctactttaatq gcacaatggattcaaatgggacatttgttaatgtaacaatgagcacatttaactggaaggatta cattggagatgacagtcacttttatgttttggatgggcaaaaagaccctttactctgtggaaat ggctcagatgcaggccagtgtccagaaggatacatctgtgtgaagqctgqtcqaaaccccaact atggctacacaagctttgacacctttagctgggctttcctgtctctatttcqactcatqactca tttgtcctggtcattttcttgggctcattttatttggtgaatttgatcctggctgtggtggcca tggcctatgaggagcagaatcaggccaccttggaagaagcagaacaaaaagaggccgaatttca gcagatgctcgaacagcttaaaaagcaacaggaagaagctcaggcagttgcggcagcatcaqct gcttcaagagatttcagtggaataggtgggttaggagagctgttggaaagttcttcagaagcat gcaccttgaaggaaacaacaaggagagagagacagctttcccaaatccgaatctgaagacagc gtcaaaagaagcagcttccttttctccatggatggaaacagactgaccagtgacaaaaaattct getececteateagtetetettgagtateegtggeteeetgtttteeeceaagaegeaatageaa aacaagcattttcagtttcagaggtcgggcaaaggatgttggatctgaaaatgactttgctgat gatgaacacagcacatttgaagacagcgaaagcaggaqacatcactqtttqtqccqcacaqac atggagagcgacgcaacagtaacgttagtcaggccagtatgtcatccaggatggtgccagggct ccttcagctctaacgtcacctactggacaacttcccccagagggcaccaccacagaaacggaaq tcagaaagagaaggttaagctcttaccagatttcaatggagatgctggaggattcctctqqaaq gcaaagagccgtgagcatagccagcattctgaccaacacaatggaagaacttgaagaatctaga cagaaatgtccgccatgctggtatagatttgccaatgtgttcttgatctgggactgctgtgatg catggttaaaagtaaaacatcttgtgaatttaattgttatggatccatttqttgatcttqccat cactatttgcattgtcttaaataccctctttatggccatggagcactaccccatgactgaqcaa ttcagtagtgttgactgtaggaaacctggtctttactgggattttcacagcagaaatqqttc tcaagatcattgccatggatccttattactatttccaagaaggctggaatatctttgatggaat tattgtcagcctcagtttaatggagcttggtctgtcaaatgtggagggattgtctgtactgcga tcattcagactgcttagagttttcaagttggcaaaatcctggcccacactaaatatqctaatta  ${\tt agatcattggcaattctgtgggggctctaggaaacctcaccttggtgttggccatcatcgtctt}$ catttttgctgtggtcggcatgcagctctttggtaagagctacaaagaatgtgtctgcaagatc 

### FIGURE 1 (continued)

tccgcgtgctgtgtggagagtggatagagaccatgtgggactgtatggaggtcgctggccaaac catgtgccttattgtttcatgttggtcatggtcattggaaaccttgtggttctgaacctcttt ctggccttattgttgagttcatttagctcagacaaccttgctgctactgatgatgacaatgaaa tgaataatctgcagattgcagtaggaagaatgcaaaagggaattgattatgtgaaaaataagat gcgggagtgtttccaaaaagccttttttagaaagccaaaagttatagaaatccatgaaggcaat aagatagacagctgcatgtccaataatactggaattgaaataagcaaagagcttaattatctta gagatgggaatggaaccaccagtggtgtaggtactggaagcagtgttgaaaaatacgtaatcga tgaaaatgattatatgtcattcataaacaaccccagcctcaccgtcacagtgccaattgctgtt ggagagtctgactttgaaaacttaaatactgaagagttcagcagtgagtcagaactagaagaaa agaaggtgaacaagctgaaactgaacccgaagaagaccttaaaccggaagcttgttttactgaa ggatgtattaaaaagtttccattctgtcaagtaagtacagaagaaggcaaagggaagatctggt ggaatcttcgaaaaacctgctacagtattgttgagcacaactggtttgagactttcattgtgtt catgatccttctcagtagtggtgcattggcctttgaagatatatacattgaacagcgaaagact tcaaatgggttgcttatggatttcaaacatatttcactaatgcctggtgctggctagatttctt gatcgttgatgtttctttggttagcctggtagccaatgctcttggctactcagaactcggtgcc atcaaatcattacggacattaagagctttaagacctctaagagccttatcccggtttgaaggca tgagggtggttgtgaatgctcttgttggagcaattccctctatcatgaatgtgctgttggtctg tgtgttaacatgacaacgggtaacatgtttgacattagtgatgttaacaatttgagtgactgtc aggctcttggcaagcaagctcggtggaaaacgtgaaagtaaactttgataatgttggcgctgg gattcacgagatgttaaacttcagcctgtatatgaagaaaatctgtacatgtatttatactttq tcatctttatcatctttgggtcattcttcactctgaatctattcattggtgtcatcatagataa cttcaaccagcagaaaaagaagtttggaggtcaagacatctttatgacagaggaacagaaaaa tattacaatgcaatgaagaaacttggatccaagaaacctcagaaacccatacctcgcccagcaa acaaattccaaggaatggtctttgattttgtaaccagacaagtctttgatatcagcatcatgat cctcatctgcctcaacatggtcaccatgatggtggaaacggatgaccagggcaaatacatgacc ctagttttgtcccggatcaacctagtgttcattgttctgttcactggagaatttgtgctgaggc tegteteecteagacactactactteactataggetggaacatetttgaetttgtggtggtgat tctctccattgtaggtatgtttctggctgagatgatagaaaagtattttgtgtcccctaccttg ttccgagtgatccgtcttgccaggattggccgaatcctacgtctgatcaaaggagcaaaggga tccgcacgctgctctttgctttgatgatgtcccttcctgcgttgtttaacatcggcctcctqct cttcctggtcatgtttatctatgccatctttgggatgtccaactttgcctatgttaaaaaqqaa gctggaattgatgacatgttcaactttgagacctttggcaacagcatgatctgcttgttccaaa ttacaacctctgctggctgggatggattgctagcacctattcttaatagtgcaccacccgactg tgaccctgacacaattcaccctggcagctcagttaagggagactgtgggaacccatctgttggg attttcttttttgtcagttacatcatcatatccttcctggttgtggtgaacatgtacatcgcgg tgagatgttctatgaggtttgggaaaagtttgatcccgatgcgacccagtttatagagttctct aaactctctgattttgcagctgccctggatcctcctcttctcatagcaaaacccaacaagtcc agettattgecatggatetgeceatggteagtggtgaceggatecaetgtettgatattttatt gacaggtttatggcatcaaacccctccaaagtctcttatgagcctattacaaccactttgaaac

## FIGURE 1 (continued)

### FIGURE 2: NaIII18 amino acid (SEQ ID NO: 2)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKEQDNDDENKPKPNSDLEAG KNLPFIYGDIPPEMVSEPLEDLDPYYINKKTFIVMNKGKAIFRFSATSALYILTPLNPVR KIAIKILVHSLFSMLIMCTILTNCVFMTLSNPPDWTKNVEYTFTGIYTFESLIKILARGF CLEDFTFLRDPWNWLDFSVIVMAYVTEFVSLGNVSALRTFRVLRALKTISVIPGLKTIVG ALIQSVKKLSDVMILTVFCLSVFALIGLQLFMGNLRNKCLQWPPSDSAFETNTTSYFNGT MDSNGTFVNVTMSTFNWKDYIGDDSHFYVLDGQKDPLLCGNGSDAGQCPEGYICVKAGRN PNYGYTSFDTFSWAFLSLFRLMTQDYWENLYQLTLRAAGKTYMIFFVLVIFLGSFYLVNL ILAVVAMAYEEQNQATLEEAEQKEAEFQQMLEQLKKQQEEAQAVAAASAASRDFSGIGGL GELLESSSEASKLSSKSAKEWRNRRKKRRRREHLEGNNKGERDSFPKSESEDSVKRSSFL FSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRRNSKTSIFSFRGRAKDVGSENDFADDEH STFEDSESRRDSLFVPHRHGERRNSNVSQASMSSRMVPGLPANGKMHSTVDCNGVVSLVG GPSALTSPTGOLPPEGTTTETEVRKRRLSSYQISMEMLEDSSGRQRAVSIASILTNTMEE LEESRQKCPPCWYRFANVFLIWDCCDAWLKVKHLVNLIVMDPFVDLAITICIVLNTLFMA MEHYPMTEQFSSVLTVGNLVFTGIFTAEMVLKIIAMDPYYYFQEGWNIFDGIIVSLSLME LGLSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVGALGNLTLVLAIIVFIFAV VGMQLFGKSYKECVCKINDDCTLPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGOT MCLIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNEMNNLQIAVGRMQKGIDYV KNKMRECFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGS SVEKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEFSSESELEESKEKLNATSS SEGSTVDVVLPREGEQAETEPEEDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKT CYSIVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTMLEYADKVFTYIFILEMLLK WVAYGFQTYFTNAWCWLDFLIVDVSLVSLVANALGYSELGAIKSLRTLRALRPLRALSRF EGMRVVVNALVGAIPSIMNVLLVCLIFWLIFSIMGVNLFAGKFYHCVNMTTGNMFDISDV NNLSDCQALGKQARWKNVKVNFDNVGAGYLALLQVATFKGWMDIMYAAVDSRDVKLQPVY EENLYMYLYFVIFIIFGSFFTLNLFIGVIIDNFNQQKKKFGGQDIFMTEEQKKYYNAMKK LGSKKPQKPIPRPANKFQGMVFDFVTRQVFDISIMILICLNMVTMMVETDDQGKYMTLVL SRINLVFIVLFTGEFVLRLVSLRHYYFTIGWNIFDFVVVILSIVGMFLAEMIEKYFVSPT LFRVIRLARIGRILRLIKGAKGIRTLLFALMMSLPALFNIGLLLFLVMFIYAIFGMSNFA YVKKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILNSAPPDCDPDTIHPGSSVK GDCGNPSVGIFFFVSYIIISFLVVVNMYIAVILENFSVATEESAEPLSEDDFEMFYEVWE KFDPDATQFIEFSKLSDFAAALDPPLLIAKPNKVQLIAMDLPMVSGDRIHCLDILFAFTK RVLGESGEMDALRIQMEDRFMASNPSKVSYEPITTTLKRKQEEVSAAIIQRNFRCYLLKO RLKNISSNYNKEAIKGRIDLPIKQDMIIDKLNGNSTPEKTDGSSSTTSPPSYDSVTKPDK EKFEKDKPEKESKGKEVRENQK

## FIGURE 3: cDNA sequence of human SCN3A of Clare et al. (SEQ ID NO: 3)

1 taccctaacc atcttggatg ctgggctttg ttatgctgta attcataagg ctctgtttta 61 tcagagatta tggagcaaga aaactgaagc caagccacat caaggtttga cagggatgag 121 atacctgtca aggattcata gtagagtggc ttactgggaa aggagcaaag aatctcttct 181 agggatattg taagaataaa tgagataatt cacagaaggg acctggagct tttccqqaaa 241 aaggtgctgt gactatctaa ggtaattcgt atgcaagaag ctacacgtaa ttaaatgtgc 301 aggatgaaaa gatggcacag gcactgttgg tacccccagg acctgaaagc ttccgccttt 361 ttactagaga atctcttgct gctatcgaaa aacgtgctgc agaagagaaa gccaagaagc 421 ccaaaaagga acaagataat gatgatgaga acaaaccaaa gccaaatagt gacttggaag 481 ctggaaagaa ccttccattt atttatggag acattcctcc agagatggtg tcagagcccc 541 tggaggacct ggatccctac tatatcaata agaaaacttt tatagtaatg aataaaggaa 601 aggcaatttt ccgattcagt gccacctctg ccttgtatat tttaactcca ctaaaccctg 661 ttaggaaaat tgctatcaag attttggtac attctttatt cagcatgctt atcatgtgca 721 ctattttgac caactgtgta tttatgacct tgagcaaccc tcctgactgg acaaagaatg 781 tagagtacac attcactgga atctatacct ttgagtcact tataaaaatc ttggcaagag 841 ggttttgctt agaagatttt acgtttcttc gtgatccatg gaactggctg gatttcagtg 901 tcattgtgat ggcgtatgta acagaatttg taagcctagg caatgtttca gcccttcgaa 961 ctttcagagt cttgagagct ctgaaaacta tttctgtaat tccaggttta aagaccattg 1021 tgggggccct gatccagtcg gtaaagaagc tttctgatgt gatgatcctg actgtgttct 1081 gtctgagcgt gtttgctctc attgggctgc agctgttcat gggcaatctg aggaataaat 1141 gtttgcagtg gcccccaagc gattctgctt ttgaaaccaa caccacttcc tactttaatg 1201 gcacaatgga ttcaaatggg acatttgtta atgtaacaat gagcacattt aactggaagg 1261 attacattgg agatgacagt cacttttatg ttttggatgg gcaaaaagac cctttactct 1321 gtggaaatgg ctcagatgca ggccagtgtc cagaaggata catctgtgtg aaggctggtc 1381 gaaaccccaa ctatggctac acaagctttg acacctttag ctgggctttc ctgtctctat 1441 ttcgactcat gactcaagac tactgggaaa atctttacca gttgacatta cgtgctgctg 1501 ggaaaacata catgatattt tttgtcctgg tcattttctt gggctcattt tatttggtga 1561 atttgatect ggetgtggtg gecatggeet atgaggagea gaateaggee acettggaag 1621 aagcagaaca aaaagaggcc gaatttcagc agatgctcga acagcttaaa aagcaacagg 1681 aagaagetea ggeagetegeg geageateag etgetteaag agattteagt ggaataggtg 1741 ggttaggaga gctgttggaa agttcttcag aagcatcaaa gttgagttcc aaaagtgcta 1801 aagaatggag gaaccgaagg aagaaaagaa gacagagaga gcaccttgaa ggaaacaaca 1861 aaggagagag agacagcttt cccaaatccg aatctgaaga cagcgtcaaa agaagcaqct 1921 tccttttctc catggatgga aacagactga ccagtgacaa aaaattctgc tccctcatc 1981 agtetetett gagtateegt ggeteeetgt ttteeceaag acgeaatage aaaacaagea 2041 ttttcagttt cagaggtcgg gcaaaggatg ttggatctga aaatgacttt gctgatgatg 2101 aacacagcac atttgaagac agcgaaagca ggagagactc actqtttqtq ccqcacagac 2161 atggagageg acgcaacagt aacggcacca ccactgaaac qqaaqtcaga aagagaaggt 2221 taagctetta ecagatttea atggagatge tggaggatte etetggaagg caaagageeg 2281 tgagcatagc cagcattctg accaacacaa tggaagaact tgaagaatct agacagaaat 2341 gtccgccatg ctggtataga tttgccaatg tgttcttgat ctgggactgc tgtgatgcat 2401 ggttaaaagt aaaacatctt gtgaatttaa ttgttatgga tccatttgtt gatcttgcca 2461 tcactatttg cattgtctta aataccctct ttatggccat ggagcactac cccatqactq 2521 agcaattcag tagtgtgttg actgtaggaa acctggtctt tactgggatt ttcacagcag 2581 aaatggttct caagatcatt gccatggatc cttattacta tttccaagaa ggctggaata 2641 totttgatgg aattattgtc agcctcagtt taatggagct tggtctgtca aatgtggagg 2701 gattgtctgt actgcgatca ttcagactgc ttagagtttt caagttggca aaatcctggc 2761 ccacactaaa tatgctaatt aagatcattg gcaattctgt gggggctcta ggaaacctca 2821 ccttggtgtt ggccatcatc gtcttcattt ttgctgtggt cggcatgcag ctctttggta 2881 agagctacaa agaatgtgtc tgcaagatca atgatgactg tacgctccca cggtgqcaca 2941 tgaacgactt cttccactcc ttcctgattg tgttccgcgt gctgtgtgga gagtggatag 3001 agaccatgtg ggactgtatg gaggtcgctg gccaaaccat gtgccttatt gttttcatgt 3061 tggtcatggt cattggaaac cttgtggttc tgaacctctt tctggcctta ttgttgagtt 3121 catttagete agacaacett getgetaetg atgatgacaa tgaaatgaat aatetgeaga

## FIGURE 3 (continued)

3181	ttgcagtagg	aagaatgcaa	aagggaattg	attatgtgaa	aaataagatg	cgggagtgtt
				ttatagaaat		
3301	acagctgcat	gtccaataat	actggaattg	aaataagcaa	agagcttaat	tatcttagag
3361	atgggaatgg	aaccaccagt	ggtgtaggta	ctggaagcag	tgttgaaaaa	tacgtaatcg
3421	atgaaaatga	ttatatgtca	ttcataaaca	accccagcct	caccgtcaca	gtgccaattg
3481	ctgttggaga	gtctgacttt	gaaaacttaa	atactgaaga	gttcagcagt	gagtcagaac
3541	tagaagaaag	caaagagaaa	ttaaatgcaa	ccagctcatc	tgaaggaagc	acagttgatg
				aaactgaacc		
				agtttccatt		
3721	aaggcaaagg	gaagatctgg	tggaatcttc	gaaaaacctg	ctacagtatt	gttgagcaca
				tccttctcag		
				tcaaaaccat		
3901	tetttaceta	tatattcatt	ctggaaatgc	ttctcaaatg	ggttgcttat	ggatttcaaa
3961	catatttcac	taatgcctgg	tgctggctag	atttcttgat	cgttgatgtt	tctttggtta
				aactcggtgc		
				cccggtttga		
				tgaatgtgct		
				tgtttgctgg		
				gtgatgttaa		
				tgaaagtaaa		
				ttaaaggctg		
				ctgtatatga		
				cattcttcac		
				agaagtttgg		
				tgaagaaact		
				aaggaatggt		
				tctgcctcaa		
				tagttttgtc		
				agctcgtctc		
				tggtgattct		
				cccctacctt		
				aaggagcaaa		
5101	ttgctttgat	gatgtccctt	cctgcgttgt	ttaacatcgg	cctcctgctc	ttcctggtca
				actttgccta		
				gcaacagcat		
				cacctattct		
5341	gtgaccctga	cacaattcac	cctggcagct	cagttaaggg	agactgtggg	aacccatctg
5401	ttgggatttt	ctttttegtc	agttacatca	tcatatcctt	cctggttgtg	gtgaacatgt
				ttgctactga		
				tttgggaaaa		
				ttgcagctgc		
5641	tagcaaaacc	caacaaagtc	cagcttattg	ccatggatct	gcccatggtc	agtggtgacc
5701	ggatccactg	tcttgatatt	ttatttgcct	ttacaaagcg	tgttttgggt	gagagtggag
5761	agatggatgc	ccttcgaata	cagatggaag	acaggtttat	ggcatcaaac	ccctccaaag
5821	tctcttatga	gcctattaca	accactttga	aacgtaaaca	agaggaggtg	tctgccgcta
5881	tcattcagcg	taatttcaga	tgttatcttt	taaagcaaag	gttaaaaaat	atatcaagta
						gacatgatta
6001	. ttgacaaact	aaatgggaac	tccactccag	aaaaaacaga	tgggagttcc	tctaccacct
				cagacaagga		
						gaaacaaaga
6181	attatctttg	tgatcaattg	tttacagcct	atgaaggtaa	agtatatgtg	tcaactggac
				tttaacaaa		
						atcaacattg
6361	. acaagaggtt	gctgtttta	. ttaccagctg	acactgctga	ggagaaaccc	aatggctacc

## FIGURE 3 (continued)

6421 tagactatag ggatagttgt gcaaagtgaa cattgtaact acaccaaaca cctttagtac 6481 agtccttgca tccattctat ttttaacttc catatctgcc atattttac aaaatttgtt 6541 ctagtgcatt tccatggtcc ccaattcata gtttattcat aatgctatgt cactatttt

## FIGURE 4: amino acid sequence of human SCN3A (SEQ ID NO: 4)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKEQDNDDENKPKPNSDLEAGKNLPFI YGDIPPEMVSEPLEDLDPYYINKKTFIVMNKGKAIFRFSATSALYILTPLNPVRKIAIKILVHS LFSMLIMCTILTNCVFMTLSNPPDWTKNVEYTFTGIYTFESLIKILARGFCLEDFTFLRDPWNW LDFSVIVMAYVTEFVSLGNVSALRTFRVLRALKTISVIPGLKTIVGALIQSVKKLSDVMILTVF CLSVFALIGLQLFMGNLRNKCLQWPPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG DDSHFYVLDGOKDPLLCGNGSDAGQCPEGYICVKAGRNPNYGYTSFDTFSWAFLSLFRLMTQDY WENLYQLTLRAAGKTYMIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQKEAEFQQM LEQLKKQQEEAQAVAAASAASRDFSGIGGLGELLESSSEASKLSSKSAKEWRNRRKKRRQREHL EGNNKGERDSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRRNSKTS IFSFRGRAKDVGSENDFADDEHSTFEDSESRRDSLFVPHRHGERRNSNGTTTETEVRKRRLSSY QISMEMLEDSSGRQRAVSIASILTNTMEELEESRQKCPPCWYRFANVFLIWDCCDAWLKVKHLV. NLIVMDPFVDLAITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFTGIFTAEMVLKIIAMDPY YYFQEGWNIFDGIIVSLSLMELGLSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVGA LGNLTLVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCTLPRWHMNDFFHSFLIVFRVLCGEWI ETMWDCMEVAGQTMCLIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNEMNNLQIAVG RMQKGIDYVKNKMRECFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDGNGTTSG VGTGSSVEKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEFSSESELEESKEKLNATS SSEGSTVDVVLPREGEQAETEPEEDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCYS IVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTMLEYADKVFTYIFILEMLLKWVAYGFQ TYFTNAWCWLDFLIVDVSLVSLVANALGYSELGAIKSLRTLRALRPLRALSRFEGMRVVVNALV GAIPSIMNVLLVCLIFWLIFSIMGVNLFAGKFYHCVNMTTGNMFDISDVNNLSDCQALGKQARW KNVKVNFDNVGAGYLALLQVATFKGWMDIMYAAVDSRDVKLQPVYEENLYMYLYFVIFIIFGSF FTLNLFIGVIIDNFNQQKKKFGGQDIFMTEEQKKYYNAMKKLGSKKPQKPIPRPANKFQGMVFD FVTRQVFDISIMILICLNMVTMMVETDDQGKYMTLVLSRINLVFIVLFTGEFVLKLVSLRHYYF TIGWNIFDFVVVILSIVGMFLAEMIEKYFVSPTLFRVIRLARIGRILRLIKGAKGIRTLLFALM MSLPALFNIGLLLFLVMF1YA1FGMSNFAYVKKEAGIDDMFNFETFGNSMICLFQITTSAGWDG LLAPILNSAPPDCDPDTIHPGSSVKGDCGNPSVGIFFFVSYIIISFLVVVNMYIAVILENFSVA TEESAEPLSEDDFEMFYEVWEKFDPDATQFIEFSKLSDFAAALDPPLLIAKPNKVQLIAMDLPM VSGDRIHCLDILFAFTKRVLGESGEMDALRIQMEDRFMASNPSKVSYEPITTTLKRKQEEVSAA IIQRNFRCYLLKQRLKNISSNYNKEAIKGRIDLPIKQDMIIDKLNGNSTPEKTDGSSSTTSPPS YDSVTKPDKEKFEKDKPEKESKGKEVRENOK

## FIGURE 5: cDNA of human sodium channel a-subunit variant by Jeong et al. (SEQ ID NO: 5)

1 agegaagegg aggeataage agagaggatt etggaaaggt etetttgttt tettateeae 61 agagaaagaa agaaaaaaaa ttgtaactaa tttgtaaacc tctqtggtca aaaaaaaaa 121 aaaaaaaaa gctgaacagc tgccagagga agacacgtta taccctaacc atcttqgatq 181 ctgggctttg ttatgctgta attcataagg ctctgtttta tcagagatta tggagcaaga 241 aaactgaagc caagccacat caaggtttga cagggatgag atacctgtca aggattcata 301 gtagagtggc ttactgggaa aggagcaaag aatctcttct agggatattg taagaataaa 361 tgagataatt cacagaaggg acctggagct tttccggaaa aaggtgctgt gactatctaa 421 ggtaattcgt atgcaagaag ctacacgtaa ttaaatgtgc aggatgaaaa gatggcacag 481 gcactgttgg tacccccagg acctgaaagc ttccgccttt ttactagaga atctcttgct 541 gctatcgaaa aacgtgctgc agaagagaaa gccaagaagc ccaaaaagga acaagataat 601 gatgatgaga acaaaccaaa gccaaatagt gacttggaag ctggaaagaa ccttccattt 661 atttatggag acattectee agagatggtg teagageece tggaggacet ggateectae 721 tatatcaata agaaaacttt tatagtaatg aataaaggaa aggcaatttt ccgattcagt 781 gccacctctg ccttgtatat tttaactcca ctaaaccctg ttaggaaaat tgctatcaag 841 attttggtac attctttatt cagcatgctt atcatgtgca ctattttgac caactgtgta 901 tttatgacct tgagcaaccc tcctgactgg acaaagaatg tagagtacac attcactgga 961 atctatacct ttgagtcact tataaaaatc ttggcaagag ggttttgctt agaagatttt 1021 acgtttcttc gtgatccatg gaactggctg gatttcagtg tcattgtgat ggcatatgtg 1081 acagagtttg tggacctggg caatgtctca gcgttgagaa cattcagagt tctccgagca 1141 ctgaaaacaa tttcagtcat tccaggttta aagaccattg tgggggccct gatccagtcg 1201 gtaaagaage tttetgatgt gatgateetg actgtgttet gtetgagegt gtttgetete 1261 attgggctgc agctgttcat gggcaatctg aggaataaat gtttgcagtq gcccccaaqc 1321 gattetgett ttgaaaccaa caccacttee taetttaatq qeacaatqqa tteaaatqqq 1381 acatttgtta atgtaacaat gagcacattt aactggaagg attacattgg agatgacagt 1441 cacttttatg ttttggatgg gcaaaaagac cctttactct gtggaaatgg ctcagatgca 1501 ggccagtgtc cagaaggata catctgtgtg aaggctggtc gaaaccccaa ctatqqctac 1561 acaagctttg acacctttag ctgggctttc ctgtctctat ttcgactcat gactcaagac 1621 tattgggaaa atctttacca qttgacatta cqtqctqctq qqaaaacata catqatattt 1681 tttgtcctgg tcattttctt gggctcattt tatttggtga atttgatcct ggctgtggtg 1741 gccatggcct atgaggagca gaatcaggcc accttggaag aagcagaaca aaaagaggcc 1801 gaatttcagc agatgctcga acagcttaaa aagcaacagg aagaagctca ggcagttgcg 1861 gcagcatcag ctgcttcaag agatttcagt ggagtaggtg ggttaggaga gctgttggaa 1921 agttetteag aageateaaa gttgagttee aaaggtgeta aagaatggag gaaceggagg 1981 aagaaaagaa gacagagaga gcaccttgaa ggaaacaaca aaggagagag agacagcttt 2041 cccaaatccg aatctgaaga cagcgtcaaa agaagcagct tccttttctc catggatgga 2101 aacagactga ccagtgacaa aaaattctgc tcccctcatc agtctctctt gagtatccgt ·2161 ggctccctgt tttccccaag acgcaatagc aaaacaagca ttttcagttt cagaggtcgg 2221 gcaaaggatg ttggatctga aaatgacttt gctgatgatg aacacagcac atttgaagac 2281 ggcgaaagca ggagagactc actgtttgtg ccgcacagac atggagagcg acgcaacagt 2341 aacgttagtc aggccagtat gtcatccagg atggtgccag ggcttccagc aaatgggaag 2401 atgcacagea ctgtggattg caatggtgtg gtttccttgg tgggtggacc ttcagctcta 2461 acgtcaccta ctggacaact tcccccagag ggcaccacca ctgaaacgga agtcagaaag 2521 agaaggttaa gctcttacca gatttcaatg gagatgctgg aggattcctc tggaaggcaa 2581 agagccgtga gcatagccag cattctgacc aacacaatgg aagaacttga agaatctaga 2641 cagaaatgtc cgccatgctg gtatagattt gccaatgtgt tcttgatctg ggactgctgt 2701 gatgcatggt taaaagtaaa acatcttgtg aatttaattg ttatggatcc atttgttgat 2761 cttgccatca ctatttgcat tgtcttaaat accetettta tggccatgga gcactacce 2821 atgactgage aatteagtag tgtgttgact gtaggaaace tggtetttae tgggatttte 2881 acagcagaaa tggttctcaa gatcattgcc atggatcctt attactattt ccaaqaaggc 2941 tygaatatet ttgatggaat tattgteage eteagtttaa tygagettgg tetgteaaat 3001 gtggagggat tgtctgtact gcgatcattc agactgctta gagttttcaa gttqqcaaaa 3061 teetggeeca cactaaatat getaattaag atcattggea attetgtggg ggetetagga 3121 aacctcacct tggtgttggc catcatcgtc ttcatttttg ctgtggtcgg catqcaqctc

## FIGURE 5 (continued)

	-				
3181 tttggtaaga	gctacaaaga	atgtgtctgc	aagatcaatg	atgactgtac	gctcccacgg
3241 tggcacatga	acgacttctt	ccactccttc	ctgattgtgt	tccgcgtgct	gtgtggagag
3301 tggatagaga	ccatqtqqqa	ctqtatqqaq	gtcgctggcc	aaaccatgtg	ccttattgtt
3361 ttcatgttgg	tcatggtcat	tggaaacctt	gtggttctga	acctctttct	ggccttatta
3421 ttgagttcat	ttagctcaga	caaccttqct	gctactgatg	atgacaatga	aatgaataat
3481 ctgcagattg					
3541 gagtgtttcc					
3601 aagatagaca					
3661 cttagagatg					
3721 gtaatcgatg	aaaatgatta	tatgtcattc	ataaacaacc	ccagcctcac	cgtcacagtg
3781 ccaattgctg	ttqqaqaqtc	tgactttgaa	aacttaaata	ctgaagagtt	cagcagtgag
3841 tcagaactag	aagaaagcaa	agagaaatta	aatgcaacca	gctcatctga	aggaagcaca
3901 gttgatgttg	ttctaccccq	agaaggtgaa	caagctgaaa	ctgaacccga	agaagacttt
3961 aaaccggaag	cttqttttac	tgaagggtgt	attaaaaaqt	ttccattctq	tcaagtaagt
4021 acagaagaag	qcaaaqqqaa	gatctggtgg	aatcttcgaa	aaacctgcta	cagtattgtt
4081 gagcacaact	ggtttgagac	tttcattqtq	ttcatqatcc	ttctcagtag	tggtgcattg
4141 gcctttgaag	atatatacat	tgaacagcga	aagactatca	aaaccatgct	agaatatgct
4201 gacaaagtct	ttacctatat	attcattctq	qaaatqcttc	tcaaatgggt	tgcttatgga
4261 tttcaaacat	atttcactaa	tacctaatac	tqqctagatt	tcttgatcgt	tgatgtttct
4321 ttggttagcc					
4381 cggacattaa					
4441 gttgtgaatg	ctcttqttqq	agcaattccc	tctatcatga	atgtgctgtt	ggtctgtctc
4501 atcttctggt	tgatctttag	catcatgggt	gtgaatttgt	ttgctggcaa	gttctaccac
4561 tgtgttaaca					
4621 tgtcaggctc					
4681 ggcgctggct					
4741 tatgcagctg					
4801 atgtatttat	actttqtcat	ctttatcatc	tttgggtcat	tcttcactct	gaatctattc
4861 attggtgtca					
4921 tttatgacag					
4981 cctcagaaac					
5041 accagacaag					
5101 atggtggaaa					
5161 gtgttcattg					
5221 tacttcacta	. taggctggaa	catctttgac	tttgtggtgg	tgattctctc	cattgtaggt
5281 atgtttctgg	ctgagatgat	agaaaagtat	tctgtgtccc	ctaccttgtt	ccgagtgatc
5341 cgtcttgcca					
5401 ctgctctttg	ctttgatgat	gtcccttcct	gcgttgttta	acatcggcct	cctgctcttc
5461 ctggtcatgt	ttatctatgc	catctttggg	atgtccaact	ttgcctatgt	taaaaaggaa
5521 gctggaattg	atgacatgtt	caactttgag	acctttggca	acagcatgat	ctgcttgttc
5581 caaattacaa	cctctgctgg	ctgggatgga	ttgctagcac	ctattcttaa	tagtgcacca
5641 cccgactgtg					
5701 ccatctgttg	ggattttctt	ttttgtcagt	tacatcatca	. tatccttcct	ggttgtggtg
5761 aacatgtaca	tcgcggtcat	cctggagaac	ttcagtgttg	ctactgaaga	aagtgcagag
5821 cccctgagtg	g aggatgactt	tgagatgttc	tatgaggttt	gggaaaagtt	tgatcccgat
5881 gcgacccagt	ttatagagtt	ctctaaactc	tctgattttg	cagctgccct	ggatcctcct
5941 cttctcatag	g caaaacccaa	caaagtccag	cttattgcca	tggatctgcc	catggtcagt
6001 ggtgaccgga	a tccactgtct	. tgatatttta	tttgccttta	. caaagcgtgt	tttgtgtgag
6061 agtggagaga	a tggatgccct	tcgaatacag	atggaagaca	ı ggtttatggo	atcaaacccc
6121 tccaaagtct					
6181 gccgctatca					
6241 tcaagtaact					
6301 atgattatt	g acaaactaaa	tgggaactco	actccagaaa	ı aaacagatg <u>c</u>	gagttcctct
. 6361 accacccct	c ctccttccta	tgatagtgta	acaaaaccag	g acaaggaaaa	gtttgagaaa

## FIGURE 5 (continued)

6	421	gacaaaccag	aaaaagaaag	caaaggaaaa	gaggtcagag	aaaatcaaaa	gtaaaaagaa
6.	481 8	acaaagaatt	atctttqtqa	tcaattqttt	acagcctatg	aaggtaaagt	atatgtgtca
6	541 a	actggacttc	aaqaqqaqqt	ccatqccaaa	ctgactgttt	taacaaatac	tcatagtcag
6	601 t	tocctataca	agacagtgaa	gtgacctctc	tgtcactgca	actctgtgaa	gcagggtatc
6	661 8	aacattaaca	agaggttgct	gtttttatta	ccagctgaca	ctgctgagga	gaaacccaat
6	721	gactacctag	actataggga	tagttgtgca	aagtgaacat	tgtaactaca	ccaaacacct
6	781	ttagtagagt	ccttqcatcc	attctatttt	taacttccat	atctgccata	tttttacaaa
6	841 8	atttqttcta	gtgcatttcc	atggtcccca	attcatagtt	tattcataat	gctatgtcac
6	901	tatttttqta	aatgaggttt	acgttgaaga	aacagtatac	aagaaccctg	tctctcaaat
6	961	gatcagacaa	aggtgttttg	ccagagagat	aaaatttttg	ctcaaaacca	gaaaaagaat
7	021	tgtaatggct	acagtttcag	ttacttccat	tttctagatg	gctttaattt	tgaaagtatt
7	081	ttagtctgtt	atgtttgttt	ctatctgaac	agttatgtgc	ctgtaaagtc	tcctctaata
7	141	tttaaaggat	tatttttatg	caaagtattc	tgtttcagca	agtgcaaatt	ttattctaag
7	201	tttcagagct	ctatatttaa	tttaggtcaa	atgctttcca	aaaagtaatc	taataaatcc
7	261	attctagaaa	aatatatcta	aagtattgct	ttagaatagt	tgttccactt	tctgctgcag
7	321	tattgctttg	ccatcttctg	ctctcagcaa	agctgatagt	ctatgtcaat	taaataccct
7	7381	atgttatgta	aatagttatt	ttatcctgtg	gtgcatgttt	gggcaaatat	atatatagcc
7	441	tgataaacaa	cttctattaa	atcaaatatg	taccacagtg	tatgtgtctt	ttgcaagctt
7	7501	ccaacaggga	tgtatcctgt	atcattcatt	aaacatagtt	taaaggctat	cactaatgca
7	7561	tattaatatt	gcctatgctg	ctctatttta	ctcaatccat	tcttcacaag	tcttggttaa
7	7621	agaatgtcac	atattggtga	tagaatgaat	tcaacctgct	ctgtccatta	tgtcaagcag
7	7681	aataatttga	agctatttac	aaacaccttt	acttttgcac	ttttaattca	acatgagtat
7	7741	catatggtat	ctctctggat	ttcaaggaaa	cacactggat	actgcctact	gacaaaacct
7	7801	attcttcata	ttttgctaaa	aatatgtcta	aaacttgttt	aaatataaat	aatgtaaaaa
7	7861	tataatcaac	tttatttgtc	agcattttgt	acataagaaa	attattttca	ggttgatgac
7	7921	atcacaattt	attttacttt	atgcttttgc	ttttgatttt	taatcacaat	tccaaacttt
-	7981	tgaatccata	agatttttca	atggataatt	tcctaaaata	aaagttagat	aatgggtttt
8	8041	atggatttct	ttgttataat	atattttcta	ccattccaat	aggagataca	ttggtcaaac
8	B101	actcaaacct	agatcatttt	ctaccaacta	tggttgcctc	aatataacct	tttattcata
8	8161	gatgttttt	tttattcaac	ttttgtagta	tttacgtatg	cagactagtc	ttatttttt
1	8221	aattcctgct	gcactaaagc	tattacaaat	ataacatgga	ctttgttctt	tttagccatg
;	8281	aacaaagtgg	caaagttgtg	caattaccta	. acatgatata	aatttttgtt	ttttgcacaa
;	8341	accaaaagtt	taatgttaat	tcttttaca	aaactattta	ctgtagtgta	ttgaagaact
	8401	gcatgcaggg	aattgctatt	. gctaaaaaga	atggtgagct	acgtcattat	tgagccaaaa
	8461	gaataaattt	catttttat	tgcatttcac	ttattgggct	ctggggtttt	ttgtttttgt
	8521	tttttgctgt	tggcagttta	ı aaatatatat	. aattaataaa	acctgtgctt	gatctgacat
	8581	ttgtatacat	aaaagtttac	: atgaatttta	ı caacaaacta	gtgcatgatt	caccaagcag
	8641	tactacagaa	caaaggcaaa	ttaaaagcag	g ctttgtgaac	ttttatgtgt	gcaaaggatc
	8701	aagttcacat	gttccaactt	tcaggtttga	taataatagt	agtaaccacc	tacaatagct
	8761	ttcaatttca	attaactcco	: ttggctataa	gcatctaaac	tcatcttctt	tcaatataat
	8821	tgatgctato	c tootaattad	: ttggtggcta	a ataaatgtta	cattctttgt	tacttaaatg
	8881	cattatata	a actcctatgt	: atacataagg	g tattaatgat	atagttattg	agaatttata
	8941	ttaactttt	tttcaagaa	c ccttggattt	atgtgaggtc	aaaaccaaac	tcttattctc
	9001	agtggaaaa	c tccagttgta	a atgcatatti	ttaaagacaa	. tttggatcta	aatatgtatt
			cccataata	a attatataa	g gtggaaaaaa	aaaaaaaaa	aaaaaaaaaa
*	9121	aaa					

## FIGURE 6: amino acid sequence of human sodium channel $\alpha$ -subunit variant by Jeong et al. (SEQ ID NO: 6)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKEQDNDDENKPKPNSDLEAGKNLPFI YGDIPPEMVSEPLEDLDPYYINKKTFIVMNKGKAIFRFSATSALYILTPLNPVRKIAIKILVHS LFSMLIMCTILTNCVFMTLSNPPDWTKNVEYTFTGIYTFESLIKILARGFCLEDFTFLRDPWNW LDFSVIVMAYVTEFVDLGNVSALRTFRVLRALKTISVIPGLKTIVGALIQSVKKLSDVMILTVF CLSVFALIGLOLFMGNLRNKCLQWPPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG DDSHFYVLDGQKDPLLCGNGSDAGQCPEGY1CVKAGRNPNYGYTSFDTFSWAFLSLFRLMTQDY WENLYQLTLRAAGKTYMIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQKEAEFQQM LEQLKKQQEEAQAVAAASAASRDFSGVGGLGELLESSSEASKLSSKGAKEWRNRRKKRRQREHL EGNNKGERDSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRRNSKTS IFSFRGRAKDVGSENDFADDEHSTFEDGESRRDSLFVPHRHGERRNSNVSQASMSSRMVPGLPA NGKMHSTVDCNGVVSLVGGPSALTSPTGQLPPEGTTTETEVRKRRLSSYQISMEMLEDSSGROR AVSIASILTNTMEELEESRQKCPPCWYRFANVFLIWDCCDAWLKVKHLVNLIVMDPFVDLAITI CIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFTGIFTAEMVLKIIAMDPYYYFQEGWNIFDGIIV SLSLMELGLSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVGALGNLTLVLAIIVFIF AVVGMQLFGKSYKECVCKINDDCTLPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNEMNNLQIAVGRMQKGIDYVKNKMRE CFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSVEKYVIDEN DYMSFINNPSLTVTVPIAVGESDFENLNTEEFSSESELEESKEKLNATSSSEGSTVDVVLPREG EQAETEPEEDFKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCYSIVEHNWFETFIVFMI LLSSGALAFEDIYIEORKTIKTMLEYADKVFTYIFILEMLLKWVAYGFQTYFTNAWCWLDFLIV DVSLVSLVANALGYSELGAIKSLRTLRALRPLRALSRFEGMRVVVNALVGAIPSIMNVLLVCLI FWLIFSIMGVNLFAGKFYHCVNMTTGNMFDISDVNNLSDCQALGKQARWKNVKVNFDNVGAGYL ALLOVATFKGWMDIMYAAVDSRDVKLOPVYEENLYMYLYFVIFIIFGSFFTLNLFIGVIIDNFN OOKKKFGGODIFMTEEOKKYYNAMKKLGSKKPOKPIPRPANKFOGMVFDFVTRQVFDISIMILI CLNMVTMMVETDDQGKYMTLVLSRINLVFIVLFTGEFVLKLVSLRHYYFTIGWNIFDFVVVILS IVGMFLAEMIEKYSVSPTLFRVIRLARIGRILRLIKGAKGIRTLLFALMMSLPALFNIGLLLFL VMFIYAIFGMSNFAYVKKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILNSAPPDCDP DTIHPGSSVKGDRGDPSVGIFFFVSYIIISFLVVVNMYIAVILENFSVATEESAEPLSEDDFEM FYEVWEKFDPDATOFIEFSKLSDFAAALDPPLLIAKPNKVOLIAMDLPMVSGDRIHCLDILFAF TKRVLCESGEMDALRIQMEDRFMASNPSKVSYEPITTTLKRKQEEVSAAIIQRNFRCYLLKQRL KNISSNYNKEAIKGRIDLPIKQDMIIDKLNGNSTPEKTDGSSSTTPPPSYDSVTKPDKEKFEKD KPEKESKGKEVRENOK

								Section 1
	(1)	1	,10	20	)	30		48
ClareAJ251507	(1)							
huNaIII18 (AK) JeongAF225987	(1)	7 G C C 7	AGCGGAGG	 			 G	
Consensus	(1)	nucun	AGCGGAGG	CAIAAGC	nonone	.02111010	GAMMOOTO	.1011101
		***						- Section 2
	(49)	49	60		,70	80		96
ClareAJ251507	(1)							
huNaIII18 (AK)	(1)							
JeongAF225987 Consensus	(49) (49)	тттст	TATCCACA	GAGAAAG	AAAGAF	TAAAAAAI	TGTAACTA	ATTTGTA
Consensus	(40)		···					- Section 3
	(97)	97		110	,120		130	144
ClareAJ251507	(1)							
huNalli18 (AK)	(1)							
JeongAF225987		AACCT	CTGTGGTC	AAAAAA	AAAAA	AAAAAAA	AGCTGAA	CAGCTGCC
Consensus	(97)							_ Section 4
	(145)	145	.150	.160		170	.180	192
ClareAJ251507	(1)		1100					SCTTTGTT
huNaIII18 (AK)	(1)							
JeongAF225987		AGAGG	GAAGACACG					
Consensus	(145)			TAC	CTAAC	CATCTTGG	ATGCTGG	GCTTTGTT — Section 5
	(400)	402	200	210		220	230	240
ClareAJ251507	(193)		200 NGTAATTCI		i i i i i i i i i i i i i i i i i i i			
huNallI18 (AK)	(1)							
JeongAF225987	(193)	ATEC	CHARDIC	TAAGGG	MAGRA	TTATCAGI	GATTATE	GAGGAAGA
Consensus	(193)	ATGCT	rgtaattc <i>i</i>	ATAAGGC'	rctgtt	TTATCAGA	GATTATG	
								— Section 6
Ol A 1054507	(241)	241	250		260	270		288
ClareAJ251507 huNaIII18 (AK)	(81)	AAAC	reavacceas	NUCCACA	e e e e e e	737.347.47	COMMEN	<b></b>
JeongAF225987			Very Telefort					
Consensus	(241)	AAAC'	TGAAGCCA	AGCCACA	<b>TCAAGG</b>	TTTGACA	GGATGAG	ATACCTGT
								Section 7
	(289)	289	30	00	310	32	0	330
ClareAJ251507	(129)	CAAG	CANVICATION	guagagy.	BGERRY	ergggaa.	NGGREGAN	eve with other
huNaIII18 (AK) JeongAF225987	(1) (290)		GATTCATA	enarazan	ecenns	Andreas	A COMPANY	ACABOTEM
Consensus			GATTCATA					
00110011000	\_00	,						

												Section
	(337)				350		360			70		38
ClareAJ251507	(177)	rocu	AGGG!	TTATT	GTAR	THAT	YPAA2	AGA	r taku	CACAC	AAG	BGACC
huNaIII18 (AK)	(1)										، سے سے ہ محمد محمد	
leongAF225987	(337)	CTCT	aggg)	ATAIT	GTAA	61-6-61	<b>NAME OF</b>	AGA'	LLKAI	CACAC	AAG	BGACC
Consensus	(337)	TTCT	AGGG	$\mathbf{rr}\mathbf{Ar}$	GTAA	GAAT	TAAA	SAGA	rtaat	CACAC		
												Section
•	(385)	385	390		400			110		420	AP CH (70	43
		Assert Parent Lat 5	CHAL	rccgc	AAAA	TODA	ecro:	DADI	TATCI	PARGO	TAA	TCGTA
huNaIII18 (AK)	(1)											
JeongAF225987	(385)	GGAG	CTTT	recge	AAAA	AGGT	<b>GCTG</b>	rgac	TATC	AAGG	TAA	negna
Consensus	(385)	GGAG	CTTT	TCCGG	BAAAA	AGGT	GCTG	rgac	TATCT	CAAGG		
											_	Section 1
	(433)	433	44			450	market A saturate	460		,47		4
	(273)	CCAA	GRING	CUAC	CGTA	KINE				DAAAA		
huNaIII18 (AK)	(1)									GAAAA		
JeongAF225987	(433)	E 16.313	GARG	CTAC	COUN	ATTA	AATO	TGCA	GGAT(	GAAAA	GATG	GCACA
Consensus	(433)	GCAA	GAAG	CTAC	ACGTA	ATTA	AATG'	TGCA	GGAT	GAAAA	GATG	GCAC
												Section 1
	(481)	481		490		500			510			5
ClareAJ251507										CGCCT		
huNaIII18 (AK)	(17)	GCAC	TGŤT	GGTA	cccc	CAGGA	CCTG	AAAG	CTTC	CGCCT'	тттт	ACTAC
JeongAF225987	(481)	GCAC	TGTT	GGTA	cccc	CAGGA	CCTG	AAAG	CTTC	CGCCT'	TTTT	ACTAC
Consensus	(481)	GCAC	TGTT	GGTA	cccc	CAGGA	CCTG	AAAG	CTTC	CGCCT		
											\$	Section '
	(529)			54			550		560			5
ClareAJ251507										GAAGA		
huNalli18 (AK)	(65)	GAA	гстст	TGCT	GCTAT	CGAA	AAAC	GTGC	TGCA	GAAGA	GAAA	GCCA
JeongAF225987	(529)	GAAS	CTCT	TGCT	GCTAT	CGAA	AAAC	GTGC	TGCA	GAAGA	GAAA	GCCA
Consensus	(529)	GAA	rctci	TGCT	GCTAT	CGAA	AAAC	GTGC	TGCA	GAAGA	GAAA	GCCA
												Section
	(577)	577			590		600	)		610		
ClareAJ251507			CCCAA	AAAG	GAAC	AAGAT	AATG	ATGA	TGAG	AACAA	ACC	AAGC
huNalli18 (AK)										AACAA		
JeongAF225987	(577	) AAG	CCCAP	AAAG	GAAC	AAGAT	PAATG	ATG	ATGAG	AACAA	ACC	AAAGC
Consensus	(577	) AAG	CCCAP	AAAG	GAAC	AAGAT	AATG	ATG	ATGAG	AACAA	ACC	AAAGC
												Section
	(625	) 625	630		,64	0		650		660		(
ClareAJ251507	(465	) AAT.	AGTG	ACTTG	GAAG	CTGG	AAAGA	ACC	TTCCA	TATTT	TTA	rggag
	inon	יח א א	* ~ ~ ~ .		CAAC	CmcG:	אארא	ACC	מיתיים ב	rattr.	יתיתאי	rggag
huNall118 (AK)	(101	) AMI	MGIGE	マケエエの	GAAG	CIGGI	JAMOL	11 1 C C				
huNaIII18 (AK) JeongAF225987	(625	) AAT	AGTG!	ACTTG ACTTG	GAAG	CTGG1	AAAGA	AACC	TTCCA	TATT.	'TTA'	rggag

									- Section 15
	(673)	673	680		690		700	,710	720
ClareAJ251507	(513)	ATTCC	TCCAGA	GATGG	TGTCAC	AGCCC	CTGGAGO	SACCTGGA	ATCCCTAC
huNalll18 (AK)	(209)	ATTCC	TCCAGA	GATGG	TGTCAC	SAGCCC	CTGGAG	SACCTGGA	ATCCCTAC
JeongAF225987	(673)	ATTCC	TCCAGA	GATGG	TGTCAC	AGCCC	CTGGAGG	GACCTGGA	ATCCCTAC
Consensus	(673)	ATTCC	TCCAGA	GATGG	TGTCAC	AGCCC	CTGGAG	BACCTGG	ATCCCTAC
									- Section 16
	(721)	721	,730	0	740		750		768
ClareAJ251507	(561)	TATAT	CAATAA	GAAAA	CTTTT	ATAGTA	ATGAATA	AAGGAAA	AGGCAATT
huNaIII18 (AK)	(257)	TATAT	CAATAA	GAAAA	СТТТТА	ATAGTA	ATGAATA	AAAGGAA	AGGCAATT
JeongAF225987	(721)	TATAT	CAATAA	GAAAA	СТТТТ	ATAGTA	ATGAATA	AAAGGAAA	AGGCAATT
Consensus	(721)	TATAT	CAATAA	GAAAA	СТТТТ	ATAGTA	ATGAAT	AAAGGAA	AGGCAATT
									- Section 17
	(769)	769		.780	7	790	800		816
ClareAJ251507			ATTCAC	TGCCA				PTAACTC(	CACTAAAC
huNaIII18 (AK)	(305)	TTCCG	ATTCAC	TGCCA	ССТСТ	CCTTG	ተልጥልጥጥ የ	TTAACTC(	CACTAAAC
JeongAF225987									CACTAAAC
Consensus									CACTAAAC
									Section 18
	(817)	817		830		840	5	350	864
ClareAJ251507			TAGGAZ		СТАТС				PATTCAGC
huNaIII18 (AK)	(353)	ССТСТ	TAGGAZ	አልልጥጥG	CTATC	α α α α α α α α α α α α α α α α α α α	ттссть	~ A TT CT T .	TATTCAGC
JeongAF225987	(817)	ССТСТ	TAGGAZ	ነ አ አ ጥ ጥ ር	CTATC	ΔΑCΑΨΨ:	TTGGTA( ጥጥሮርጥል	CATICII	TATTCAGC
Consensus									PATTCAGC
								CHIICII.	Section 19
	(865)	865	870	, a	80	.890		900	912
ClareAJ251507									TGACCTTG
huNallI18 (AK)	(401)	ATGCT	$\mathbf{T}\mathbf{A}\mathbf{I}\mathbf{C}\mathbf{A}\mathbf{I}$	r G I G C A	יים איניטי. ריים איניטי	THE ACC	AACIGI	SIATITA.	TGACCTTG
JeongAF225987		ATGCT	татса.	ranaca ranaca	CIXII	TTGACC.	AACIGI AACTGT	3 T V T T T Y	TGACCTTG
Consensus	(865)	ATGCT	ጥልጥሮል፣	ramac a	CTATI	TTGACC.	AACIGI AACTGT	2421117	TGACCTTG
	(000)	711001	TAICA.	GIGCE	CIALL	I I GACC.	AACIGI	SIAIIIA	Section 20
	(913)	013	920		,930		940	.950	960
ClareAJ251507				TMC A CM					TCACTGGA
huNalli18 (AK)		AGCAA	CCCTC		GGACA.	A A C A A M	GTAGAG.	TACACAT.	TCACTGGA
JeongAF225987									TCACTGGA
_									TCACTGGA
Conscisus	(310)	AGCAA	CCCIC	JIGACI	GGACA.	AAGAAT	GTAGAG.	TACACAT	— Section 21
	(961)	064	97		000		000		
Clare A 1954507					,980		990		1008
ClareAJ251507 huNaIII18 (AK)									GGTTTTGC
JeongAF225987	•								GGTTTTGC
_									GGTTTTGC
Consensus	(901)	ATCTA	TACCT"	r r G A G 1	CACTT.	A'I'AAAA	ATCTTG	GCAAGAG	GGTTTTGC

						— Section 22
(1009)	1009	,1020	··	1030	_,1040	1056
ClareAJ251507 (849)	TTAGA.	AGATTTTAC	GTTTCTT	CGTGATC	CATGGAACTGG	CTGGATTTC
huNaIII18 (AK) (545)	TTAGA.	AGATTTTAC	GTTTCTI	CGTGATC	CATGGAACTGG	CTGGATTTC
JeongAF225987 (1009)	TTAGA.	AGATTTTAC	GTTTCTI	CGTGATC	CATGGAACTGG	CTGGATTTC
Consensus (1009)	TTAGA	AGATTTTAC	GTTTCTT	CGTGATC	CATGGAACTGG	CTGGATTTC
						Section 23
(1057)	1057		070	,1080	,1090	1104
ClareAJ251507 (897)	AGTGT	CATTGTGAT	GGCGTAT	GTRACAG	AATTTGTAAGC	CTAGGCAAT
huNaIII18 (AK) (593)	AGTGT	CATTGTGAT	GGCGTAT	GTÄACAG	AATTTGTAAGO	CTAGGCAAT
JeongAF225987 (1057)	AGTGT	CATTGTGAT	GGCATAT	GTGACAG	AGTTTGTGGAC	CTGGGCAAT
Consensus (1057)	AGTGT	CATTGTGAT	GGCGTAT	GTAACAG	AATTTGTAAGO	CTAGGCAAT
						Section 24
		1110	_1120	1130		1152
ClareAJ251507 (945)	GTTTC	AGCCCTTCG	AACTTT	AGAGT	TENGAGCTCTC	BAAAACKATT
huNallI18 (AK) (641)	) СТИТС	AGC CT VCG	AACTTT	CAGAGTO	TGAGAGCTCTC	SAAAACTATT
JeongAF225987 (1105)	GTCTC	AGCGTTGAG	AACATTO	CAGAGTTC	TCCGAGCACTO	SAAAACAATT
Consensus (1105)	GTTTC	AGCCCTTCG	AACTTTC	CAGAGTCT	TGAGAGCTCTC	SAAAACTATT
		<del></del>				Section 25
(1153	1153	1160	,1170	,1	180 ,11	90 1200
ClareAJ251507 (993	TCTGT	ATTCCAGG	CARATTE	SACCATTG	TGGGGGCCCTC	SATCCAGTCG
huNall118 (AK) (689	) TCEGT	ATTCCAGG	CAAATT	GACCATTG	TGGGGGCCCT	GATCCAGTCG
JeongAF225987 (1153)	TCAGT	CATTCCAGG	CAAATT	GACCATTG	TGGGGGCCCT	GATCCAGTCG
Consensus (1153	) TCTGT	AATTCCAGG	CAAATT	GACCATTG	TGGGGGCCCT	GATCCAGTCG
						Section 26
	1201	,1210	,12		,1230	1248
ClareAJ251507 (1041	) GTAAA	GAAGCTTTC	TGATGT	GATGATCC	TGACTGTGTT	CTGTCTGAGC
huNaIII18 (AK) (737						
JeongAF225987 (1201	) GTAAA	GAAGCTTTC	TGATGT	GATGATCC	TGACTGTGTT	CTGTCTGAGC
Consensus (1201	) GTAAA	GAAGCTTTC	TGATGT	GATGATCO	TGACTGTGTT	CTGTCTGAGC
						Section 27
	) 1249	,1260		,1270	,1280	1296
ClareAJ251507 (1089	) GTGTT	TGCTCTCAT	TGGGCT	GCAGCTGT	TCATGGGCAA	TCTGAGGAAT
huNall118 (AK) (785	) GTGTI	TGCTCTCAT	TGGGCT	GCAGCTGI	TCATGGGCAA	TCTGAGGAAT
JeongAF225987 (1249	) GTGTI	TGCTCTCAT	TGGGCT	GCAGCTGI	TCATGGGCAA	<b>PCTGAGGAAT</b>
Consensus (1249	) GTGTI	TGCTCTCAT	TGGGCT	GCAGCTGI	TCATGGGCAA	<b>PCTGAGGAAT</b>
						Section 28
	) 1297		310	,1320	,1330	1344
ClareAJ251507 (1137						
huNalli18 (AK) (833	) AAAT	STTTGCAGT	GCCCCC.	AAGCGATI	CTGCTTTTGA.	AACCAACACC
JeongAF225987 (1297	AAATO	STTTGCAGT	GCCCCC.	AAGCGATT	CTGCTTTTGA.	AACCAACACC
Consensus (1297	) AAATG	TTTGCAGT	GCCCCC.	AAGCGATT	CTGCTTTTGA.	AACCAACACC

		<del></del>											Section	n 29
(1	345)	1345 ,1	350		136	30		1370	)		1380			1392
ClareAJ251507 (1	185)	ACTTC	CTAC	ATTT	ATGG	CACA	DTA.	GATT	CAA	ATGG	GACA	$TT_{L}$	TGTT	$\overline{TAP}$
huNall118 (AK) (	(881)	ACTTC	CTAC	TTTA	ATGG	CACA	ATG	GATI	CAA	ATGG	GACA	TT	TGTT	TAP
JeongAF225987 (1	345)	ACTTC	CTAÇ	ATTT	ATGG	CACA	ATG	GATT	CAA	ATGG	GAC	TTL	TGTT	TAA
Consensus (1	345)	ACTTC	CTAC	TTTA	ATGG	CACA	ATG	GATI	CAA	ATGG	GAC	TT	TGTT	TAA
													Section	
	1393)		,140			1410		,1	420		,14			1440
ClareAJ251507 (1	233)	GTAAC	AATG	AGCA	CATT	TAAC	TGG	AAGG	TTA	ACAT	TGGI	AGA	TGAC	AGT
huNallI18 (AK) (	(929)	GTAAC	AATG	AGCA	CATI	TAAC	TGG	AAGG	ATT	ACAT	TGG	AGA	TGAC	AGT
JeongAF225987 (1	(393	GTAAC	AATG	AGCA	CATT	TAAC	TGG	AAGG	ATT	ACAT	TGG	AGA	TGAC	AGT
Consensus (1	(393	GTAAC	AATG	AGCA	CATT	TAAC	TGG	AAGG	ATT	ACAT	TGG	AGA	TGAC	AGT
·													Section	
(1	1441)	1441		,1450		.14	60		,147	0				1488
ClareAJ251507 (1	(1281	CACTT	ГТАТ	GTTT	TGGA	TGGG	CAA	AAAG			ACTO	TG.		
huNalII18 (AK) `(	(977)	CACTT	ГТАТ	GTTT	TGGA	TGG	CAA	AAAC	ACC	СТТТ	ACTO	TTG	TGGA	AAT
JeongAF225987 (1	(441)	CACTT	ТАТ	GTTT	TGGA	TGGG	CAA	AAAG	ACC	Cuuu	A C T C	יייה	TGGA	ጥልA
Consensus (1	1441)	CACTT	гатт	GTTT	TGGA	TGGG	CAA	AAAG	ACC	CTTT	ACTO	ንጥG	TGGA	AΑΨ
													Section	
(1	1489)	1489		,150	20		,1510			1520				1536
ClareAJ251507 (1											CTG	rem		
huNaill18 (AK) (1	1025)	GGCTC	TASA	'GCAG	CCC	CTCI		GAAC	CAT	ACAT	CTG	r G r	CAAC	CCT
JeongAF225987 (1	1489)	GGCTC	АСРТ	GCAG	GCCZ	CTCI	ירכם.	GAAC	CAT	ACAT	יכיים.	r G T	CAAC	CCT
Consensus (1	1489)	GGCTC	AGAT	GCAG	GCCZ	GTGT	ירכא	GAAG	CAT	A C A T	יכיים.	ram	CAAC	COT
										110111	C 1 G .		Section	
(1	1537)	1537			,1550		15	560		,15	70			1584
ClareAJ251507 (1			AAAC			TGGG	7 A Tr	ACAZ	GCT	TTGA	CAC	ىلىك ب		
huNalli18 (AK) (1	1073)	GGTCG	AAAC		аста	. тсс.	יש ארי	ACAZ	CCT	ጥጥር እ	CAC	~ Tr T	TAGC	ጥርር
JeongAF225987 (1	1537)	GGTCG.	AAAC	$CCC^{2}$	A C T Z	.тсс(	יש ארי	ACAZ	CCT	TTCZ	CAC	- ተ ተ	TAGC	ጥርር
Consensus (1	1537)	GGTCG	AAAC	CCCA	አ ር ጥ ጀ	ጥርርር	ים גריי	ACAZ	1001 1001	ተ ተ ር 2	CAC	ጎ ጉ ጉ	TAGC	ጥሮር
								210212	1001	1102	-		Sectio	
(1	1585)	1585	1590		,16	00		.1610	<u> </u>		.1620		COOLIO	1632
ClareAJ251507 (1	1425)	CCTTT		T C T C			CTC			77707		S T C	CCAA	
huNaIII18 (AK) (1	1121)	CCTTT	CCTC	יייכונ	ים עם עם עם עדי דידידים	rmcG2	$^{2}C$ $^{2}C$	ATC	$^{\prime}$	AAGE	mmy STY	E T G	CCAA	y y w
JeongAF225987 (1	1585)	GCTTT	CCTC	יייכייכ	י ת ע עה מ	rmcG2	$V \subset T \subset T$	ATC	7 C T C	AAGE	TITA	DUC STG	CCNN	y y u
Consensus (1	1585)	CCTTT	CCTC	3 T C T C	ית עת מייי	nmcc:	7 C T C	A TO 2		AMGE	、Cun y v	C II G	CCAA	у у ш Ч Ч Т
	1000,			,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	.1211	LICGZ	AC I C	MIGI	4C1C	MAGE	CIA		Sectio	
13	1633)	1633	,16	40		,1650			1660		16	570		1680
ClareAJ251507 (1					CIM		no om			7 7 7 7				
huNall118 (AK) (	1160\	CHUMBA	CCAC	2 1. 1. G /	CAT	TACG:	r GCT	COM	3 G G A	) AAA.	ATA	CAT	GATA	T.T.T.
JeongAF225987 (*	1100) 1622\	CTTTA	CCAC	2 T T G F	ACAT'	LACG'	recr	GCJ.(	2007 2007	) AAA	ATA	CAT	GATA	TTT
•														
Consensus (	1033)	CTTTA	CCAC	2 T. T. C. F	ACAT".	rACG'	rGCT	GC.I.(	⊈ىئىنى	AAAC	ATA	CAT	GATA	ттт

							_				900	ction 36
	(1681)	1681		,1690		.17	00		,1710			1728
ClareAJ251507	(1521)	TTTG	TCCT	GGTC	TTTA	TCTT	GGCI	CATT	ית את תיתי	ጉጥርርጥር	ጥጥልል	TGATC
nunaiii 18 (AK)	(1217)	TTTC	TCCI	GGTC	A ጥጥጥ	<b>ጥርጥጥ</b> (	$GGC^{\eta}$	$\Gamma \cap \Delta \cap \Gamma$	ኮ ጥ ላ ጥ ጥ	maama	יחידות מי	שת א שיכי
JeongAF225987	(1681)	TTTG	TCCI	GGTC	ТТТА	TCTTC	GGCT	ГСАТТ	υπΑπη	. T 00 T C	2 2 2 mm	mcnmc
Consensus	(1681)	TTTG	TCCI	GGTC	ATTT	$\mathbf{T}\mathbf{C}\mathbf{T}\mathbf{T}\mathbf{C}$	GGCT	יים איז זיים איז	ւաւև Σաևաև	ነጥርርጥር	ישטע ע עיק יישטע ע עיקי	ከርአ ሞር 1 GZ I C
												ction 37
<b>~</b> 1 <b></b>	(1729)	1729		.17	40		,1750		,1760	)		_ 1776
ClareAJ251507	(1569)	CTGG	CTGI	G.GTG	GCCA	TGGC	TATO	GAGGA	GCAGA	ATCAG	GCCA	CCTTG
nuivain 18 (AK)	(1265)	CTGG	GCTGT	GGTG	GCCA'	$\mathtt{TGGC}($	CTATO	GAGGA	GCAGA	ATCAC	GCCA	CCTTG
JeongAF225987	(1729)	CTGG	CTGI	GGTG	GCCA'	TGGC	TATO	SAGGA	GCAGA	ATCAC	GCCA	CCTTG
Consensus	(1729)	CTGG	CTGI	GGTG	GCCA	TGGC	TATO	GAGGA	GCAGA	ATCAG	GCCA	CCTTG
											_	ction 38
	(1777)	1777			,1790		,18	00		810		1824
ClareAJ251507	(1617)	GAAG	AAGC	AGAA	CAAA.	AAGA	GCCG	TTAAE	TCAGO	CAGATO	CTCG	AACAG
nuivaiii 18 (AK)	(1313)	GAAG	BAAGC	CAGAA	CAAA.	AAGAG	GCCC	FTAAE	TCAGO	CAGATO	CTCG	AACAG
JeongAF225987	(1777)	GAAG	AAGC	CAGAA	CAAA.	AAGA	GCCC	FAATI	TCAGO	AGATO	CTCG	AACAG
Consensus	(1777)	GAAG	SAAGO	AGAA	CAAA.	AAGA	GCCC	TTAAE	TCAGO	AGATO	CTCG	AACAG
												ction 39
	(1825)	1825	_,1830		.18	340	_	,1850		1860		1872
ClareAJ251507	(1665)	CTTP	AAAA	GCAA	CAGG.	AAGA	AGCTO	CAGGC	AGTTO	CGGCA	GCAT	CAGCT
nuNaiii18 (AK)	(1361)	CTTA	AAAA	GCAA	CAGG.	AAGA	AGCTC	CAGGC	AGTTO	CGGCA	GCAT	CAGCT
JeongAF225987	(1825)	CTTA	AAAA	GCAA	CAGG.	AAGA	AGCTO	CAGGC	AGTTO	CGGCA	GCAT	CAGCT
Consensus	(1825)	CTTA	AAAA	GCAA	CAGG.	AAGA	AGCTO	CAGGC	AGTTO	CGGCA	GCAT	CAGCT
												ction 40
	(1873)	1873		880		_1890		,190	00	,19	10	1920
ClareAJ251507	(1713)	GCTI	CAAC	AGAT	TTCA	GTGG	ATAC	GTGG	GTTAC	GAGAG	CTGT	TGGAA
nunaiii18 (AK)	(1409)	GCTI	CAAC	BAGAT	TTCA	GTGGZ	$\mathbf{AT}\mathbf{A}$	GTGG	GTTAG	GAGAG	CTGT	TGGAA
JeongAF225987	(1873)	GCTI	CAAG	AGAT	TTCA	GTGG	AGTAC	GTGG	GTTAG	GAGAG	CTGT	TGGAA
Consensus	(1873)	GCTI	CAAG	AGAT	TTCA	GTGG	ATA	GTGG	GTTAG	GAGAG	CTGT	TGGAA
									·			ction 41
	(1921)	1921		,1930		,19	40		1950			1968
ClareAJ251507	(1761)	AGTI	CTTC	AGAA	GCAT	CAAA	TTGA	AGTTC	CAAA	GTGCT	DAAG	AATGG
huNall118 (AK)	(1457)	AGTI	CTTC	AGAA	GCAT	CAAA	STTGA	ነር ፓርር አርጥጥር	CAAA	CTCCT	ם מ מ מי	9 9 T C C
JeongAF225987	(1921)	AGTI	CTTC	AGAA	GCAT	CAAA	STTGA	AGTTC	CAAAC	GTGCT	ים א א מ	A A TICC
Consensus	(1921)	AGTT	CTTC	AGAA	GCAT	CAAA	TTTG Z	ኒርጥጥር	CAAAA	CTCCT	AAAG.	V V A LIGG
	<u> </u>									101001	_	ction 42
	(1969)	1969		,19	980		1990		2000	)		2016
ClareAJ251507	(1809)	AGGA	ACCG	AAGG	AAGA	AAAG	AAGAC	GAG	AGAGO	, <b>y</b> C C m u	CAAC	CAAAC
huNalli18 (AK)	(1505)	AGGA	ACCC	AAGG	AAGA	AAAG	AAGAC	CGGAC	SAGAGO	, y C C u u	CAAC	CXXXC
JeongAF225987	(1969)	AGGA	ACC	GAGG	AAGA	AAAG	AAGAC	E GAG	AGAGG	CACCTI	CAMC	GAAAC
Consensus	(1969)	AGGA	ACCG	AAGG	AAGA	AAAC	ADGDO	CACAC	MONGO	. A C C I I	CAAG	CAAAC
	/						TAGA	CACAC	DOMOA	-MCCTT	GAAG	GAAAC

												Section	43
	(2017)				2030		204			2050			2064
ClareAJ251507	(1857)	AACA	AAGGA	GAGA	GAGA	CAGC	TTTC	CCAA	ATCC	GAAT	CTGA	AGACA	AGC
huNaIII18 (AK)	(1553)	AACA	AAGGA	AGAGA	GAGA	CAGC	TTTC	CCAA	ATCC	GAAT	CTGA	AGAC	AGC
JeongAF225987	(2017)	AACA	AAGGA	AGAGA	GAGA	CAGC	TTTC	CCAA	ATCO	GAAT	CTGA	AGAC	AGC.
Consensus	(2017)	AACA	AAGGA	GAGA	GAGA	CAGC	тттс	CCAP	ATCO	GAAT	CTGA	AGAC	4GC
												- Section	1 44
ClareAJ251507	(2065)	2065	2070		208	80		2090		210	0		2112
ClareAJ251507	(1905)	GTCA	AAAGA	AAGCA	GCTT	CCTT	ттст	CCAT	rggan	GGAA	ACAG	ACTG	ACC
huNaIII18 (AK)	(1601)	GTCA	AAAGA	AAGCA	GCTI	CCTT	TTCI	CCA	rggan	rggaa	ACAG	SACTG	ACC
JeongAF225987	(2065)	GTCA	AAAGA	AAGCA	GCTI	CCTT	TTCI	CCAT	rggan	rggaa	ACAG	ACTG	ACC
Consensus													
												- Section	า 45
	(2113)	2113	21	20		2130		21	40		2150		2160
ClareAJ251507			ACAA	AAAA			CCTC	CATC	AGTC	CTCT	TGAC	STATC	CGT
huNaIII18 (AK)													
												STATC	
Consensus													
												_ Section	
	(2161)	2161		2170		21	80		2190				2208
ClareAJ251507			CCCT		rccc			AATA			GCA	гтттс	AGT
huNall118 (AK)													
JeongAF225987												TTTTC	
Consensus													
												_ Sectio	
	(2209)	2209		22	220		2230		22	240			2256
ClareAJ251507						AGGA	rgrr	GGAT			ACT	TTGCT	GAT
huNaIII18 (AK)													
												TTGCT	
Consensus													
												_ Sectio	
	(2257)	2257			2270		22	280		2290			2304
ClareAJ251507			BAACA	CAGC					AAAG	CAGGA	GAG	ACTCA	CTG
huNall118 (AK													
JeongAF225987												ACTCA	
Consensus													
												_ Section	
	(2305	2305	2310		23	320		2330		23	40		2352
ClareAJ251507	7 (2145	TTT	GTGCC	GCAC	AGAC	ATGG	AGAG	CGAC	GCAA	CAGT	AACG		
huNallI18 (AK	) (1841	) TTT	GTGCC	GCAC	AGAC	ATGG	AGAG	CGAC	GCAA	CAGT	AACG	THACT	CATE
JeongAF225987												TUAGI	
Consensu													
	`												

							Section 50
	(2353)		2360	2370	2380	2390	240
ClareAJ251507							
huNallI18 (AK)	(1889)	GCCAGI	ATGTCATC	CAGGATGGI	CCAGGGCT!	<b>PCCAGCAAA</b>	Teggaa
JeongAF225987	(2353)	GCCAGI	ATGTCATC	CAGGATGGI	GCCAGGGCT	<b>PCCAGCAAA</b>	TI G G G AVAN
Consensus	(2353)	GCCAGT	ATGTCATC	CAGGATGGT	GCCAGGGCT	TCCAGCAAA	TGGGAA
		···					Section 5
	(2401)	2401	2410	2420	2430		244
ClareAJ251507	(2185)						
huNall118 (AK)	(1937)	ATGCAC	AGCACTGT	GGATTGCAP	TGGTGTGGT'	rtcerreer	GGGTGG
JeongAF225987	(2401)	ATGCAC	AGCACTGT	GGATTGCA	TGGTGTGGT'	rrcorregr	ссепас
Consensus	(2401)	ATGCAC	AGCACTGT	GGATTGCA	TGGTGTGGT	ттссттест	GGGTGG
							Section 52
	(2449)	2449	2460	24	70 24	80	249
ClareAJ251507							GCAC
			Program XX	CHESTONS.	TI GGACA'ACT!	n <i>oordina</i>	GCAC
JeongAF225987	(2449)	e e mino	remema a c		TGGACAACT		CCAC
•	(2449)	ССППС	A CCUCUDA CCUCUDA	CTCACCTAC	TGGACAACT	TCCCCCACA	CCCCAC
	(2140)		-	GICACCIAC	1 GGACAAC I	ICCCCCAGA	Section 5
	(2497)	2407	2	510	2500	000	
Clara 1251507	(2481)	2491	<del></del>	310	2520	2530	254
01d16AJZ31307	(2022)	ACCAC	GAAACGGA	AGTCAGAAA	GAGAAGGTT.	AAGCTCTTA	CCAGAT
JeongAF225987	(2407)	ACCACA	AGAAACGGA	AGTCAGAAA	AGAGAAGGTT	AAGCTCTTA	CCAGAT
	(2497)	ACCAC	gGAAACGGA	AGTCAGAAA	AGAGAAGGTT.	AAGCTCTTA	CCAGAT
Consensus	(2497)	ACCAC.	rGAAACGGA	AGTCAGAAA	AGAGAAGGTT.		
	·	0045					Section 5
01 1105/505	(2545)		2550	2560	2570	2580	259
ClareAJ251507	(2238)	TCAATO	GGAGATGCI	GGAGGATT	CCTCTGGAAG	GCAAAGAGC	CGTGAG
huNaIII18 (AK)	(2081)	TCAATO	GGAGATGCI	GGAGGATT	CCTCTGGAAG	GCAAAGAGC	CGTGAG
JeongAF225987	(2545)	TCAATO	GGAGATGCI	GGAGGATT	CCTCTGGAAG	GCAAAGAGC	CGTGAG
Consensus	(2545)	TCAAT	GAGATGCT	GGAGGATT	CCTCTGGAAG	GCAAAGAGC	CGTGAG
	<del></del>						Section 5
	(2593)		2600	2610	2620	2630	264
ClareAJ251507	(2286)	ATAGC	CAGCATTCI	GACCAACA	CAATGGAAGA	ACTTGAAGA	ATCTAG
huNaIII18 (AK)	(2129)	ATAGC	CAGCATTCT	GACCAACA	CAATGGAAGA	ACTTGAAGA	ATCTAG
JeongAF225987	(2593)	ATAGC	CAGCATTCT	GACCAACA	CAATGGAAGA	ACTTGAAGA	ATCTAG
Consensus	(2593)	ATAGC	CAGCATTCI	GACCAACA	CAATGGAAGA	ACTTGAAGA	ATCTAG
							- Section 5
	(2641)	2641	2650	2660	2670		268
ClareAJ251507	(2334)	CAGAA	ATGTCCGCC	ATGCTGGT	ATAGATTTGC	СААТСТСТТ	- <u>- 200</u>
huNall118 (AK)	(2177	CAGAA	ATGTCCGCC	CATGCTGGT	ATAGATTTGC	CAATGTGTT	C
JeongAF225987	(2641)	CAGAA	$\mathtt{ATGTCCGCC}$	CATGCTGGT	ATAGATTTGC	CAATGTGTT	ᠬᢗᠳᡎᢗ᠌᠌ᢧᡎ

									Section 57
	(2689)			2700	27		2720		2736
ClareAJ251507									
huNaIII18 (AK)	(2225)	TGGGA	CTGCTG	rgargc	ATGGT	TAAAAG'	TAAAACA	TCTTGT	GAATTTA
JeongAF225987	(2689)	TGGGA	CTGCTG!	TGATGC	ATGGT	raaaag:	TAAAACA	TCTTGT	GAATTTA
Consensus	(2689)	TGGGA	CTGCTG	TGATGC	ATGGT	TAAAAG'	TAAAACA	TCTTGT	GAATTTA
									Section 58
ClareAJ251507	(2737)	2737		2750		2760	277	0	2784
ClareAJ251507	(2430)	ATTGT	TATGGA'	TCCATI	TGTTG	ATCTTG	CCATCAC	OTTTAT	CATTGTC
huNall118 (AK)	(2273)	ATTGT	'ATGGA	TCCATT	TGTTG	ATCTTG	CCATCAC	OTTTAT	CATTGTC
JeongAF225987	(2737)	ATTGT	ratgga'	TCCATI	TGTTG	ATCTTG	CCATCAC	OTTTAT	CATTGTC
Consensus	(2737)	ATTGT	TATGGA'	TCCATI	TGTTG	ATCTTG	CCATCAC	TATTTO	CATTGTC
	·								Section 59
	(2785)	2785	2790	28		2810		2820	2832
ClareAJ251507	(2478)	TTAAA	TACCCT	CTTTAT	GGCCA	TGGAGC	ACTACCC	CATGAC	TGAGCAA
huNaIII18 (AK)	(2321)	TTAAA	TACCCT	CTTTAI	GGCCA	TGGAGC	ACTACCC	CATGAC	TGAGCAA
JeongAF225987									
									CTGAGCAA
									- Section 60
	(2833)	2833	2840		2850	28	360	2870	2880
ClareAJ251507	(2526)	TTCAG	TAGTGT	GTTGAC	TGTAG	GAAACC	тестстт	TACTGO	GATTTTC
huNall118 (AK)									
									GATTTTC
									GATTTTC
	(								Section 61
	(2881)	2881	289	90	2900		2910		2928
ClareAJ251507								TCCTTA	TTACTAT
huNaIII18 (AK)									
JeongAF225987									
									ATTACTAT
	\								- Section 62
•	(2929)	2929		2940	2	950	2960		2976
ClareAJ251507								TGTCAC	CCTCAGT
huNallI18 (AK)									
JeongAF225987									GCTCAGT
									GCTCAGT
									- Section 63
	(2977)	2977		2990		3000	30	10	3024
ClareAJ251507	(2670)	TTAAT	GGAGCT		TGTCAA				TACTGCGA
huNallI18 (AK									
JeongAF225987									TACTGCGA
									TACTGCGA
		,							

														- Section	
ClareAJ251507	(3025)	3025	303	0		3040	)		,3050			3060	)		3072
ClareAJ251507	(2718)	TCAT	TCA	GACT	GCT	ΓAGA	GTT	TTC	AAGT	TGG	CAA	$\nabla TAP$	CTG	GCCC	CACA
huNall118 (AK)	(2561)	TCAT	TCA	GACT	GCT	raga	GTT	TTC	AAGT	TGG	CAA	OTAP	CTG	GCCC	CACA
JeongAF225987	(3025)	TCAT	TCA	GACT	GCT	raga	GTT	TTCI	AAGT	TGG	CAA	$\Delta TC$	CTC	GCCC	CACA
Consensus	(3025)	TCAT	TCA	GACT	GCT	raga	GTT	TTC	AAGT	TGG	CAA	OTAA	CTC	GCCC	CACA
														- Section	on 65
	(3073)			3080			090		3	100			110		3120
ClareAJ251507															
huNallI18 (AK)															
	(3073)														
Consensus	(3073)	CTAA	ATA	$\mathtt{TGCI}$	TAA	PAAT	SATC	TTA	GGCA	ATT	'CTG'	rgge	GGG	CTCTI	AGGA
														<ul> <li>Section</li> </ul>	on 66
	(3121)			31:	30		314			,318					3168
ClareAJ251507	(2814)	AACC	TCA	CCTI	GGT	GTTC	GCC	ATC	ATCG	TCT	TCA	TTTI	TGC	TGT	GTC
huNall118 (AK)															
JeongAF225987															
Consensus	(3121)	AACC	CTCA	CCTI	GGT	GTT	GCC	ATC	ATCG	TCI	TCA	TTT	rtgo	CTGT	GGTC
															on 67
	(3169)	3169			3180			3190			3200				3216
ClareAJ251507	(2862)	GGC	TGC	AGCT	CTT	TGG	raag	AGC'	TACA	AAG	AAT	GTG	rcro	GCAA	GATC
huNall118 (AK)	(2705)	GGC	ATGC	AGCI	CTT	TGG	raag	AGC'	TACA	AAG	TAA	GTG	гсто	GCAA	GATC
JeongAF225987	(3169)	GGC	ATGC	AGCI	CTT	TGG	raag	AGC'	TACA	AAG	TAAG	GTG	гсто	GCAA	GATC
Consensus	(3169)	GGC	ATGC	AGCT	CTT	TGG	raag	AGC	TACA	AAG	TAAG	GTG	rcro	GCAA	GATC
														_ Secti	
	(3217)	3217			32	230		,32	240		.32	250			3264
ClareAJ251507	(2910)	AATO	SATG	ACTO	TAC	GCT	CCCA	CGG	TGGC	ACA	TGA	ACG	ACT	CTT	CCAC
huNall118 (AK															
JeongAF225987	(3217)	AATO	GATG	ACTO	STAC	GCT	CCCA	CGG	TGGC	AC	TGA	ACG	ACT	CTT	CCAC
Consensus															
															ion 69
	(3265)	3265	327	70		328	0		3290	)		330	0		3312
ClareAJ251507	7 (2958)	TCC	TTCC	TGA	TTGT	GTT	CCGC	GTG	CTGT	GTC	GAG	AGT	GGA'	TAGA	GACC
huNall118 (AK	) (2801)	TCC	TTCC	TGA	гтст	GTT	CCGC	GTG	CTGT	GTO	GAG	AGT	GGA	TAGA	GACC
JeongAF225987	(3265)	TCC	TTCC	TGAT	TTGT	GTT	ccgc	GTG	CTGT	GTC	GAG	AGT	GGA	TAGA	GACC
Consensu															
														- Sect	
	(3313)	3313		3320			3330		3	340			3350		3360
ClareAJ25150	7 (3006	ATG	TGGG	АСТ	GTAT	GGA	GGTC	CGCT	GGCC	:AA	ACCA	TGT	GCC	ጥጥ A ጥ	ጥርጥባ
huNall118 (AK															
JeongAF225987		) ATG													
Consensu															
		,													

													- Section	on 71
	(3361)	3361		3370	)		,3380	)		3390				3408
ClareAJ251507	(3054)	TTCAT	GTT	GGTC	ATG	GTC	TTC	GAAA	ACCT	TGT	GGTTC	TGA.	ACCTO	TTT
huNallI18 (AK)	(2897)	TTCAT	GTT	GGTC	ATG	STCA	ATTO	GAAA	CCT	ጥርጥ	GGTTC	ጥርል	A C C T C	արարար
JeongAF225987	(3361)	TTCAT	GTT	GGTC	ATG	STCA	$\Sigma TT$	GGAA	ACCT	TGT	GGTTC	TGA	ACCTC	ጥጥጥ
Consensus	(3361)	TTCAT	GTT	GGTC	ATG	GTC	ATTC	GAAA	ACCT	TGT	GGTTC	TGA	ACCTO	TTT
													- Section	
	(3409)			,3	420		,3	3430		.34	440			3456
ClareAJ251507	(3102)	CTGGC	CTT	ATT	TTG	AGT	rca1	ATTT	SCTC	AGA	CAACC	TTG	CTGCT	ACT
huNaIII18 (AK)	(2945)	CTGGC	CTT	ATTÇ	TTG	AGT	rcan	OATTI	GCTC	AGA	CAACC	TTG	CTGCT	ACT
JeongAF225987	(3409)	CTGGC	CTT	ATTA	TTG	AGT:	rcan	OATTI	GCTC	AGA	CAACC	TTG	CTGCT	ACT
Consensus	(3409)	CTGGC	CTT	ATTO	TTG	AGT	rcan	CTTAC	GCTC	AGA	CAACC	TTG	CTGCT	АСТ
													- Section	
	(3457)	3457			347	0		3480	С		3490			3504
ClareAJ251507	(3150)	GATGA	TGA	CAAI	GAA	ATG	AATA	AATCI	rgca	GAT	TGCAG	TAG	GAAGA	ATG
huNaIII18 (AK)	(2993)	GATGA	TGA	CAAT	GAA	ATGA	ATA	AATCI	rgca	GAT	TGCAG	TAG	GAAGA	ATG
JeongAF225987	(3457)	GATGA	TGA	CAAT	GAA	ATG	AATA	AATCI	rgca	GAT	TGCAG	TAG	GAAGA	ATG
Consensus	(3457)	GATGA	TGA	CAAT	GAA	ATG	AATA	AATCI	rgca	GAT	TGCAG	TAG	GAAGA	AΤG
													- Section	
	(3505)	3505	3510		;	3520		3	3530		35	40		3552
ClareAJ251507	(3198)	CAAAA	GGG	TTAA	TAP	T A T	STG A	KAAA	ααπα	CAT	GCGGG	10 C T	CTTTC	2002
huNaIII18 (AK)	(3041)	CAAAA	GGG	רתאא	TATT	ኮልጥ(	3461	XAAA	ממים	ת עט.	60666	2001	CTTTC	יר א א
JeongAF225987	(3505)	CAAAA	GGG	rraa	ית ב בינ	יייביי	3 T G 2	X	ממחים	ע ע ט	GCGGG	NO T	COMMO	ת תיחים
Consensus	(3505)	CAAAA	GGG	דידי ב ב	יתאטי	ኮልጥ(	ኃጥር 2	2 Z Z Z Z	ልጥ ል ል	ርልጥ	GCGGG	NO I	COUNTRO	יר א א
									1 4 111	OAI		TOA.	Section	
	(3553)	3553	3	560		35	70		358	30		3590		3600
ClareAJ251507	(3246)	AAAGO	CTT	ጥጥጥባ	AGA	AAGO	CA	AAAGT	יים איים	AGA	AATICC	A THE	N N C C C	יח ג גי
huNaIII18 (AK)	(3089)	AAAGO	· Cጥጥ	4444	AGA	AAGO		AAAGI	ኮጥጥጥ	תטת מטמי	$\Delta \Delta T C C$	יא תים. מת גי	AAGGC	ית אינ יחא אי
JeongAF225987	(3553)	AAAGC	יריתיי	ጥጥጥባ	AGA	AAGO			ኮጥልጣ	אטא אטמי	$\Delta \Delta T C C$	. אדט מחגי	AAGGC	ישעע.
Consensus	(3553)	AAAGO	ירייטי	<b>ሲ</b> ተተተተ	AGA	AAG	~ C A 2	מממ	ר בע ד	תםת גם גי	N $N$ $T$ $C$ $C$	יא יים	AACCC	, y y w
								inno	LTVI	AGA	MAICC	.AIG	— Section	
	(3601)	3601		3610	1		362	n		3630			0600	3648
ClareAJ251507			1 A C A			N m C r	TCC	N 7N 17N 7N 7	N M N C	2020	7 7 mmc	1777	m > > 0 c	3040
huNall118 (AK)	(3137)	AACAT	מטת. מטמי	CAGO	TUGC.	א שרכי		v v u v v	7 T. 73 C	TGG	AATT	AAA	TAAG(	AAA
JeongAF225987	(3601)	AAGAT	. בסב יאכא	CAGC	TUCC	M T G .		y y m y z	7 T. A.C.	TGG	AATT	AAA	TAAGU	CAAA
Consensus	(3601)	AACAT	מטמי מטמי	CAGO	ים פרי. משפרי	M T G .	T C C Z	7 7 M 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	JATAC	.TGG	AATTO	AAA	TAAGO	CAAA
	(0001)	AAGAI	ngn	CAGC	, I GC.	are.	1002	AN T.W.	ATAC	.166	AATT	AAA		
	(3640)	3640		2	660			2070			000		Section	
ClareAJ251507	(3649) (3342)	CA C C C	101 7 7	3	OOU	202		3670		3	680			3696
hullalli40 / A L/	(334 <i>2)</i> (340 <i>E</i> )	CACCI	.T.A.A.	TTAT	CTT.	AGA	GAT'	3 G G A A	ATGG	AAC	CACCA	GTG	GTGT	AGGT
huNall118 (AK) JeongAF225987	(3640) (3100)	CACCE	TAA	L'A'I.	CTT.	AGA(	GAT(	GGA <i>l</i>	ATGG	AAC	CACCA	GTG	GTGT	AGGT
	(3640)	GAGCT	TAA	TTAT	rctt.	AGA	GAT'	3GGA	ATGG	AAC	CACCA	GTG	GTGT	AGGT
Consensus	(3049)	GAGCI	.TAA	.T.T.W.]	rCTT.	AGA(	) 'I'A ک	€GA <i>l</i>	ATGG	AAC	CACC	GTG	GTGT	AGGT

				Section 78
(3697) 3697	3710	3720	3730	3744
ClareAJ251507 (3390) ACTGGAAGCAGTG	GTTGAAAA	ATACGTAATC	GATGAAAATG	ATTATATG
huNall118 (AK) (3233) ACTGGAAGCAGTG	GTTGAAAA	ATACGTAATC	GATGAAAATG	ATTATATG
JeongAF225987 (3697) ACTGGAAGCAGTG	GTTGAAAA	ATACGTAATO	GATGAAAATG	ATTATATG
Consensus (3697) ACTGGAAGCAGTG	GTTGAAAA	ATACGTAATC	GATGAAAATG	ATTATATG
				- Section 79
(3745) 3745 3750	3760	3770	3780	3792
ClareAJ251507 (3438) TCATTCATAAACA	AACCCCAG	CCTCACCGT	CACAGTGCCAA	TTGCTGTT
huNall118 (AK) (3281) TCATTCATAAACA	AACCCCAG	CCTCACCGT	CACAGTGCCAA	TTGCTGTT
JeongAF225987 (3745) TCATTCATAAACA	AACCCCAG	CCTCACCGTO	CACAGTGCCAA	TTGCTGTT
Consensus (3745) TCATTCATAAACA	AACCCCAG	CCTCACCGT	CACAGTGCCAA	TTGCTGTT
<u> </u>				- Section 80
( <b>3793</b> ) 3793 ,3800	3810	,3820		
ClareAJ251507 (3486) GGAGAGTCTGACT	TTTGAAAA	CTTAAATACT	GAAGAGTTCA	GCAGTGAG
huNall118 (AK) (3329) GGAGAGTCTGACT	TTTGAAAA	CTTAAATACT	rgaagagttca	GCAGTGAG
JeongAF225987 (3793) GGAGAGTCTGACT	TTTGAAAA	CTTAAATAC	rgaagagttca	GCAGTGAG
Consensus (3793) GGAGAGTCTGACT	TTTGAAAA	CTTAAATAC	rgaagagttca	GCAGTGAG
				- Section 81
(3841) 3841 ,3850	38	360	3870	3888
ClareAJ251507 (3534) TCAGAACTAGAAC	GAAAGCAA	AGAGAAATT	AAATGCAACCA	GCTCATCT
huNall118 (AK) (3377) TCAGAACTAGAAC				
JeongAF225987 (3841) TCAGAACTAGAAC				
Consensus (3841) TCAGAACTAGAAG	GAAAGCAA	AGAGAAATT	AAATGCAACCA	GCTCATCT
				- Section 82
(3889) 3889 39	900	3910	3920	3936
ClareAJ251507 (3582) GAAGGAAGCACA	GTTGATGT'	TGTTCTACC	CCGAGAAGGTG	AACAAGCT
huNalli18 (AK) (3425) GAAGGAAGCACA	GTTGATGT	TGTTCTACC	CCGAGAAGGTG	AACAAGCT
JeongAF225987 (3889) GAAGGAAGCACA	GTTGATGT	TGTTCTACC	CCGAGAAGGTG	AACAAGCT
Consensus (3889) GAAGGAAGCACA				
				Section 83
(3937) 3937	3950	3960	3970	3984
ClareAJ251507 (3630) GAAACTGAACCC	GAAGAAGA	CETTAAACC	GGAAGCTTGTT	TTACTGAA
huNall118 (AK) (3473) GAAACTGAACCC	GAAGAAGA	COTTAAACC	GGAAGCTTGTT	TTACTGAA
JeongAF225987 (3937) GAAACTGAACCC	GAAGAAGA	CTTTAAACC	GGAAGCTTGTT	TTACTGAA
Consensus (3937) GAAACTGAACCC	GAAGAAGA	CCTTAAACC	GGAAGCTTGTT	TTACTGAA
	<del></del>			Section 84
(3985) 3985 3990	4000	,4010	4020	4032
ClareAJ251507 (3678) GG TGTATTAAA	AAGTTTCC	ATTCTGTCA.	AGTAAGTACAG	
huNall118 (AK) (3521) GG TGTATTAAA	AAGTTTCC	ATTCTGTCA	AGTAAGTACAG	SAAGAAGGC
JeongAF225987 (3985) GGGTGTATTAAA	AAGTTTCC	ATTCTGTCA	AGTAAGTACAG	GAAGAAGGC
Consensus (3985) GGATGTATTAAA				

							Secti	ion 85
ClareAJ251507	(4033)	4033	4040	,4050	,40	060	4070	4080
ClareAJ251507	(3726)	AAAGGG	AAGATCTG	GTGGAAT	CTTCGAA	AAACCTGCT	ACAGTAT	TGTT
nuNaIII18 (AK)	(3569)	AAAGGG	AAGATCTG	GTGGAAT	CTTCGAA	AAACCTGCT	TACAGTAT	TGTT
JeongAF225987	(4033)	AAAGGG	AAGATCTG	GTGGAAT	CTTCGAA	AAACCTGCT	PACAGTAT	ጥርጥጥ
Consensus	(4033)	AAAGGG	AAGATCTG	GTGGAAT	CTTCGAA	AAACCTGCT	PACAGTAT	тстт
					<del></del>	<del></del>	Sect	ion 86
	(4081)	4081	<u>,4090</u>	<u>,410</u>	0	,4110		4128
ClareAJ251507	(3774)	GAGCAC	AACTGGTT	TGAGACT	TTCATTG	TGTTCATGA	TCCTTCT	CAGT
huNalli18 (AK)	(3617)	GAGCAC	AACTGGTI	TGAGACT	TTCATTG	TGTTCATGA	ATCCTTCT	CAGT
JeongAF225987	(4081)	GAGCAC	AACTGGTT	TGAGACT	TTCATTG	TGTTCATGA	ATCCTTCT	CAGT
Consensus	(4081)	GAGCAC	AACTGGTT	TGAGACT	TTCATTG	TGTTCATGA	ATCCTTCT	CAGT
	·						Sect	
	(4129)	4129	,4140	) ,	4150	4160		4176
ClareAJ251507	(3822)	AGTGGT	GCATTGGC	CTTTGAA	GATATAT	ACATTGAAC	CAGCGAAA	GACT
huNaIII18 (AK)	(3665)	AGTGGT	GCATTGGC	CTTTGAA	GATATAT	ACATTGAAC	CAGCGAAA	GACT
JeongAF225987	(4129)	AGTGGT	GCATTGGC	CTTTGAA	GATATAT	ACATTGAAC	CAGCGAAA	GACT
Consensus	(4129)	AGTGGT	GCATTGGC	CTTTGAA	GATATAT	ACATTGAA	CAGCGAAA	GACT
								ion 88
	(4177)	4177	4	190	4200	4210		4224
ClareAJ251507	(3870)	ATCAAA	ACCATGCT	AGAATAT	GCTGACA	AAGTCTTT	A C C T A T A T	A TUTO
huNaIII18 (AK)	(3713)	ATCAAA	ACCATGCT	TAGAATAT	GCTGACA	2 2 CT CT T T		አጥጥሮ
JeongAF225987	(4177)	ATCAAA	ACCATGCT	TAGAATAT	GCTGACA	AAGTCTTT		<b>አጥጥ</b> ር
	(4177)	ATCAAA	ACCATGCT	AGAATAT	GCTGACA	AAGTCTTT	$^{1}$	ል ጥ ጥ C
								ion 89
	(4225)	4225 4	230 ·	4240	4250	42	260	4272
ClareAJ251507	(3918)	ATTCTC		ית כיתים א א	#200	CULTATION	DUU CAAAC	4212
huNaIII18 (AK)	(3761)	ልጥጥርጥር		א א מיתיריתים איז.	TGGGTIG	CTIAIGGA.	THUMCAAAC	WI WI
JeongAF225987	(4225)	ATTCTC	GAAATGC1		TGGGTTG	CTTATGGA:	LTICAAAC	WI WI
	(4225)	ATTCTC	GAAATGC1	ייירית מא א	TGGGT TG	CTTATGGA	THE CAMAC	WI WI
	(			CICAAA	1000110	CITALGGA.	Sect	
	(4273)	1273	4280	4290	A	300		
Clare A 1254507	(4213) (3066)	4213	4200	4290		300	4310	4320
ClareAJ251507	(0000)	TTCACT	AATGCCTC	GTGCTGG	CTAGATT	TCTTGATC	GTTGATGT	TTCT
huNallI18 (AK) JeongAF225987	(3003) ( (4272)	TTCACT	AATGCCTC	GTGCTGG	CTAGATT	TCTTGATC	GTTGATGT	TTCI
~	(4213) (4272)	TTCACT	AATGCCTC	GTGCTGG	CTAGATT	TCTTGATC	GTTGATGT	TTCT
Consensus	6 (4213)	TTCACT	AATGCCTC	GTGCTGG	CTAGATT	TCTTGATC		
							Seci	tion 91
01	(4321)		4330	434		4350	,	4368
ClareAJ251507	(4014)	TTGGTT	AGCCTGG	RAGCCAAT	GCTCTTG	GCTACTCA	GAACTCGG	TGCC
huNalli18 (AK)	) (3857)	TTGGT	AGCCTGG	PAGCCAAT	GCTCTTG	GCTACTCA	GAACTCGG	TGCC
JeongAF225987	(4321)	TTGGTT	PAGCCTGG	TAGCCAAT	GCTCTTG	GCTACTCA	GAACTCGG	TGCC
Consensus	s (4321)	TTGGT	AGCCTGG	PAGCCAAT	GCTCTTG	GCTACTCA	${ t GAACTCGG}$	TGCC

											Section	า 92
	(4369)	4369		4380	)	,4:	390	,4	1400			4416
ClareAJ251507	(4062)	ATCAA	ATCAT	TACO	GACA	TTAA	GAGCT	TTAAC	SACCTC'	raag	AGCC'	ATT
huNaIII18 (AK)	(3905)	ATCAA	ATCAT	TACG	GACA	TTAA	GAGCT	TTAAC	SACCTC'	TAAG	AGCC'	TTA
JeongAF225987	(4369)	ATCAA	ATCAT	TACG	GACA	ТТАА	GAGCT	TTAAC	SACCTC'	TAAG	AGCC	ATT
Consensus	(4369)	ATCAA	ATCAT	TAC	GACA	TTAA	GAGCT	TTAA	SACCTC	TAAG	AGCC	TTA
											Section	
	(4417)	4417		.4	430		4440		4450			4464
ClareAJ251507			GTTTC	GAAGO	CATG	AGGG		GTGA	ATGCTC	TTGT	TGGA	GCA
huNaIII18 (AK)	(3953)	TCCCG	GTTT	GAAGO	CATG	AGGG	TGGTT	GTGA	ATGCTC	TTGT	TGGA	GCA
JeongAF225987	(4417)	TCCCG	GTTT	GAAGC	CATG	AGGG	тсстт	GTGA	ATGCTC	 ጥጥGጥ	тGGA	GCA
Consensus												
											- Sectio	
	(4465)	4465	4470		4480		.449	90	450	0		4512
ClareAJ251507				ATCAT								
huNalli18 (AK)												
JeongAF225987	(4465)	ATTCC	ירייריי	מייר אי	TGAAT	GTGC	ጥርጥጥር	CTCT	GTCTCA	T	CTCC	ጥጥር
Consensus	(4465)	ATTCC	CTCT	атса:	rgaat	GTGC	ጥርጥጥር	CTCT	3TCTCI	ተርተነ ጥሮጥባ	CTCC	ጥጥር
							.10110	0101	JICICH		- Sectio	
	(4513)	4513	452	O	4	530		4540		4550		4560
ClareAJ251507	(4206)	ATCTI	77 A G C	<u> </u>	recen.	GTG A	V drade	mmmc	CTCCCA	N C TTT		
huNaIII18 (AK)	(4049)	ጀጥርጥባ 221 C T 3	ישאַכר:	ימ חית מ	recen	なってい	አጥጥጥር አ	ጥጥጥር	CTGGCA CTGGCA	አርጥባ ሊርጉኒ	ירתאר מעשטי	CAC
JeongAF225987									CTGGCA			
Consensus												
	(	11101	11100.		10001	0102		,1110	CIGGCA	AGII	- Sectio	
	(4561)	4561		4570		4580	)	459	n		- 000110	4608
ClareAJ251507					C A A C C					GTG 7	m c m m	
huNallI18 (AK)	(4204) (4097)	TGTGT	. בממכי	A T GAY		COTI	$\mathcal{L}$	ישמים. ישמים	7 C Y W W Y	CTC	, 27 G T T	AAC
JeongAF225987	(4561) (4561)	ጥርጥርባ	ייי א א כי	አጥርአ እጥርአ	CAACC	CCTZ	$\lambda \Delta C \Delta T C$	emmmc	ACATTA ACATTA	CTCI	, 40, 40, 41	אאכ
Consensus												
Oonschaa	(4001)	1010.	I IAAC.	AIGA	CAACC	GGIA	MCMIC	1110	ACALIA		- Sectio	
	(4609)	4600		462	n		630		4640			4656
ClareAJ251507			n C 2 C M							COMO		
huNalil18 (AK												
JeongAF225987									AAGCTC			
Consensus												
Consensu	5 (4005)	AAII.	IGAGI	GACI	GICAC	age I.	- 1 1 G G C	AAGC	AAGCTC	GGT	– Sectio	
	/40573	4057	· · · · · · · · · · · · · · · · · · ·		4070		4000		4000		- Secuc	
Ol A 105450	(4657)				4670		4680		,4690			4704
ClareAJ25150												
huNallI18 (AK												
JeongAF225987									GCTATC			
Consensu	s (4657 <sub>)</sub>	) GTGA	AAGTA	AACT	TTGA	TAAT	STTGG	CGCTG	GCTATC	TTG	CACTO	CTT

															on 99
ClareAJ251507	(4705)	4705	4710	)		4720			4730			4740			4752
ClareAJ251507	(4398)	CAAG	TGGC	CAC	PTTA	AAA	GGCT	'GGA'	TGG	r A T P	TAT	GTA	rgc	CAGC	rgtt
huNaIII18 (AK)	(4241)	CAAG	TGGC	CAC	PTTA	'AAA'	GGCT	'GGA'	TGG	r A T A	TAT	GTA	TGC	CAGC	rgtt
JeongAF225987	(4705)	CAAG	TGG	CAC	PTTA	'AAA'	GGCT	'GGA'	TGG	ATA	TAT	GTA	TGC	CAGC	rgtt
Consensus	(4705)	CAAG	TGGC	CCAC	$\Gamma T T A$	'AAA'	GGCT	'GGA'	TGG	ra T.A	TAT	GTA	TGC	CAGC	rgtt
														Sectio	n 100
		4753		760		.47	70			08		47	790		4800
ClareAJ251507	(4446)	GATT	CAC	BAGA	TGTT	AAA	CTTC	AGC	CTG	PATA	ATGA	AGA	AAZ	TCT	GTAC
huNall118 (AK)	(4289)	GATT	CAC	SAGA	TGTI	AAA	CTTC	AGC	CTG	rat <i>i</i>	ATGA	AGA	AAA	ATCT	GTAC
JeongAF225987	(4753)	GATT	CAC	SAGA	TGTI	AAA	CTTC	AGC	CTG	TAT!	ATGA	AGA	AAA	ATCT	GTAC
Consensus	(4753)	GATT	CAC	GAGA	TGTI	AAA	CTTC	AGC	CTG	TAT!	ATGA	AGA	AAA	ATCT	GTAC
												<del>,</del>		Sectio	
	(4801)	4801		481	10		4820	)		4830	C				4848
ClareAJ251507	(4494)	ATGT	TTTA	ATA	CTTI	GTC	ATCI	ATT	TCA	гст	rTGG	GTC	ATI	CTT	CACT
huNaIII18 (AK)	(4337)	ATGT	TTTA	ATA	СТТЭ	GTC	ATCI	ATT	TCA!	rcr	rTGG	GTC	ΑΤЭ	гстт	CACT
JeongAF225987	(4801)	ATGT	ATT	ATA	CTTI	GTC	ATCI	ATT	TCA	rcri	rrge	GTC	ATT	CTT	CACT
Consensus	(4801)	ATGT	ATT	ATA	CTTI	GTC	ATCI	ATT	TCA	rcT7	rrge	GTC	AT?	СТТ	CACT
														Section	
	(4849)	4849			4860		4	870		.4	1880				4896
ClareAJ251507			ATC			CGT	GTCA	TCA	TAG			CAA	CCZ	AGCA	GAAA
huNaIII18 (AK)	(4385)	CTGA	ATC	TTAT	CATT	CGT	GTCA	TCA	TAG	ATA	ACTI	CAA	CCZ	AGCA	GAAA
JeongAF225987	(4849)	CTGA	ATC	TTAT	CATT	rggr	GTCA	TCA	TAG	ATA	ACTI	CAA	CCZ	AGCA	GAAA
Consensus	(4849)	CTGA	ATC	TTAT	CATT	rggr	GTCA	TCA	TAG	ATA	ACTI	CAA	CCZ	AGCA	GAAA
	·													Section	
	(4897)	4897			,49	10		492	20		49	30			4944
ClareAJ251507	(4590)	AAGA	AGT	rTGG	AGG	CAA	GACA	TCT	TTA	TGA	CAGA	GGA	ACA	AGAA	AAAA
huNaIII18 (AK)	(4433)	AAGA	AGT	rrgg	AGG	CAA	GACA	TCT	TTA	TGA	CAGA	GGA	AC	AGAA	AAAA
JeongAF225987	(4897)	AAGA	AGT	rrgg	AGG	CAA	GACA	TCT	TTA	TGA	CAGA	GGA	ACA	AGAA	AAAA
Consensus	(4897)	AAGA	AGT	rrgg	AGG	CAA	GACA	TCT	TTA	TGA	CAGA	GGA	AC	AGAA	AAAA
														Section	
	(4945)	4945	495	0		4960			4970			4980			4992
ClareAJ251507	(4638)	TATT	ACA	ATGC	AATO	GAAG	AAAC	TTG	GAT	CCA	AGAZ	ACC	TC	AGAA	ACCC
huNaIII18 (AK)	(4481)	TTAT	ACA.	ODTA	TAA	GAAG	AAAC	TTG	GAT	CCA	AGA <i>I</i>	AACC	TCI	AGAA	ACCC
JeongAF225987	(4945)	TTAT	ACA	ATGO	TAA	GAAG.	AAAC	TTG	GAT	CCA	AGA	AACC	TCI	AGAA	ACCC
Consensus	(4945)	TATT	ACA	ATGC	TAA	GAAG.	AAAC	TTG	GAT	CCA	AGA	ACC	TC	AGAA	ACCC
	<u>`                                    </u>														n 105
	(4993)	4993	,	5000		.50	010		50	020		5	030		5040
ClareAJ251507					AGC	AAAC	AAAT	TCC	AAG	GAA	TGGT			ት ጥጥጥ A ጥጥጥ	
huNaIII18 (AK)	(4529)	ATAC	CTC	GCCC	CAGC	AAAC	AAAT	TTCC	AAG	GAA	TGGT	СТТ	TG	ጥጥጥ	ТСТА
JeongAF225987	(4993)	ATAC	CTC	GCCC	CAGC	AAAC	AAA	rTCC	AAG	GAA	TGGT	СТТ	TG.	תייים אינים	TGTA
Consensus	(4993)	ATAC	CTC	GCCC	CAGC	AAAC	AAAT	rTCC	AAG	GAA	TGGT	СТТ	TG	ውጥጥ ጉጉጉ	ТСТА
	/							0							

													Section	106
	(5041)	5041		,505	0		5060	)	·	5070				5088
ClareAJ251507	(4734)	ACCA	GACA	AGT	СТТІ	GAT	ATCA	GCA:	CAT	GATC	CTCA	TCTC	GCCTC	AAC
huNallI18 (AK)	(4577)	ACCA	GACA	AGT	CTTI	GAT	ATCA	GCA	CAT	GATC	CTCA	тстс	SCCTC	CAAC
JeongAF225987	(5041)	ACCA	GACA	AGT	CTTI	GAT	ATCA	GCA	гсат	GATC	CTCA	тсто	SCCTO	CAAC
Consensus	(5041)	ACCA	GACA	AGT	СТТТ	GAT	ATCA	GCA	CAT	GATC	CTCA	тст	CCTC	CAAC
													Section	
	(5089)	5089			5100		,5	110		512	20			5136
ClareAJ251507	(4782)	ATGG	TCAC	CAT	GATO	GTG	GAAA	CGG	ATGA	CCAG	GGCA	ААТ	ACATO	20 A C C
huNall118 (AK)	(4625)	ATGG	TCAC	CAT	GATO	GTG	GAAA	CGG	ATGA	CCAG	GGCA	מתממ	$\Delta C \Delta T C$	SACC
JeongAF225987	(5089)	ATGG	TCAC	CAT	GATO	GTG	GAAA	CGG	ATGA	CCAG	GGCA	ልልጥ?	$\Delta C \Delta T C$	2200
Consensus	(5089)	ATGG	TCAC	CAT	GATO	GTG	GAAZ	CGG	АТСА	CCAC	GGCA	አልጥ:	$\Delta C \Delta T C$	2000
													Section	
	(5137)	5137			51	50		516	0					
ClareAJ251507	(4830)	CTAG	TTTT	GTC	CCGG	ATC	AACC	TAG	TGTT	CATI	GTTC	TGT	TCACT	rGGA
huNaIII18 (AK)	(4673)	CTAG	тттт	GTC	CCGG	SATC	AACC	TAG	TGTT	CATI	GTTC	TGT	TCAC	rgga
JeongAF225987	(5137)	CTAG	TTTI	GTC	CCGC	SATC	AACC	TAG	TGTT	CATT	GTTC	тст	TCAC	rgga
Consensus	(5137)	CTAG	тттт	GTC	CCGG	ATC	AACC	TAG	TGTT	САТТ	GጥጥC	ጥርጥ	TCAC	rgga
	·												Sectio	
	(5185)	5185	5190	)		5200	•		5210		522			
ClareAJ251507	(4878)	GAAT	TTGT	GCT	GAÃC	CTC	GTET	rccc	TCAG	ACAC	ТАСТ	ACT	TCACI	בסבט
huNaIII18 (AK)	(4721)	GAAT	TTGT	GCT	GAG	CTC	GT	rccc	TCAG	ACAC	ТАСТ	ים כתי	TCACT	מחמי
JeongAF225987	(5185)	GAAT	TTGI	GCT	GAÃ	CTC	GTTT	CCC	TCAG	ACAC	ТАСТ	יאכיזי	TCAC	מיתא
Consensus	(5185)	GAAT	TTGI	GCT	GAAC	CTC	GTCT	rccc	TCAG	ACAC	ТАСТ	'ACጥ'	TCAC	מיתמי
													Sectio	
	(5233)	5233	,5	240		,5	250		526	0		5270		5280
ClareAJ251507	(4926)	GGCT	GGAZ	CAT	CTTT	rgac	TTTC	TGG	TGGT	GATT	CTCT	CCA	ттст	AGGT
huNallI18 (AK)	(4769)	GGCT	GGA	CAT	CTTT	rgac	тттс	TGG	TGGT	GATT	יכידכיד	CCA	ጥጥርጥ	AGGT
JeongAF225987	(5233)	GGCT	GGA	CAT	CTTT	rgac	TTTC	TGG	$_{ m TGGT}$	GATT	СТСТ	CCA	ጥጥርጥ <i>!</i>	1001 466T
Consensus	(5233)	GGCT	GGA	ACAT	CTTT	rgac	TTTC	TGG	ጥGGጥ	GATT	יכייכיי	CCA	ጥጥርጥ	A G G T
													Sectio	
	(5281)	5281		529	90		5300	)		5310				5328
ClareAJ251507	(4974)	ATGT	TTCT	rGGC	TGAC	SATO	ATAC	AAA	ATDA	முரு நீர	יפייפיי	CCC	CTACI	TTTG
huNaIII18 (AK)	(4817)	ATGT	TTCI	rggc	TGAC	OT A F	ATA	AAA	аста	արարարարարարարարարարարարարարարարարարար	າດຫດາ		CTMC	crrc
JeongAF225987	(5281)	ATGT	TTCI	rggc	TGAG	SATO	AΤΑ	AAA	аста	اما شار شار در معالم است	າດຫວາ		CTMC	CTTO
Consensus	(5281)	ATGT	TTCI	rggc	TGA	3 A T C	ATA	AAA	AGTA	արարարա 1 1 2 2 3	CTCT		CTAC	CIIG
											. 0101		- Sectio	
	(5329)	5329			5340			5350		53	60		- 00000	5376
ClareAJ251507	(5020)	TTCC	GAGI	r G A T	1000	س کیا	1600°	AGGA	ጥጥርር	CCG	ATCC	ית א רי	C T C T	2310
huNaIII18 (AK)	(4865)	יייייייייייייייייייייייייייייייייייייי	GAGI	rao. Tabu	יררפי	$\Gamma \subset \Gamma \subset \Gamma$	GCC2	A C C A	ጠውሮር	CCC	/ Y W C C	ישע כי	GICT.	CATC
JeongAF225987	(5329)	TTCC	GAG	י פעז דעטי	יבכט. יבכני	$\Gamma \subset \Gamma$	GCC2	A C C A	JUT C C	CCC	7 M M C C	TAC	CMCM	GATC
Consensus	(5320)	1 T C C	GAG.	ועטע רפעז		1.C T 7	1000	A C C A	# 1 G G	CCGF	2 W T C C	TAC	GTCT	GATC
00119011908	(OUES)	1100	JAG:	LGWI	. CCG	$\tau \cap TA$	. GCCI	A G G A	1166	CCGA	$^{7}$ W.T.C. $^{\prime}$	TAC	GTCT	GATC

															– Se	ection	113
(	(5377)	5377				5390	)		54	00		,5	410				5424
ClareAJ251507	(5070)	AAAG	GAG	CAA	AGG	GGZ	ATC	CGC	ACG	CTGC	TC	гттс	GCT'	TTG?	TG	ATG	rcc
huNaIII18 (AK)	(4913)	AAAG	GAG	CAA	AGG	GGA	ATC	CGC	ACG	CTGC	TC	TTTC	GCT!	TTGA	ΛTG	ATG	rcc
JeongAF225987	(5377)	AAAG	GAG	CAA	AGG	GGA	ATC	CGC	ACG	CTG	TC	TTTC	GCT'	TTG	чTG	ATG	TCC
Consensus																	
																ection	
ClareAJ251507 huNallI18 (AK)	(5425)	5425	54	30		Ę	5440			5450	)		54	160		:	5472
ClareAJ251507	(5118)	CTTC	CTC	CGT	TGT	TTT	AAC	ATC	GGC	CTC	CTG	CTCT	TTC	CTGC	3TC	ATG	TTT
huNalli18 (AK)	(4961)	CTTC	CTO	CGI	TGT	TTT	AAC	ATC	GGC	CTC	CTG	CTC	rTC	CTG	3TC	ATG	${f T}{f T}{f T}$
JeongAF225987	(5425)	CTTC	CTC	GCGI	TGT	TTT	AAC	ATC	GGC	CTC	CTG	CTC	TTC	CTG	3TC	'ATG'	TTT
Consensus																	
																ection	
	(5473)	5473		5480	)		54	90			5500			551			
ClareAJ251507	(5166)	ATC	CATO	GCCF	ATCI	rTT	GGG	ATG	TCC.	AAC	TTT	GCC:	TAT	GTT	AAA	AAG	GAA
huNaIII18 (AK)	(5009)	ATC	TATO	GCCZ	ATCI	ртт	GGG	ATG	TCC.	AAC	ттт	GCC	TAT	GTT	AAA	AAG	GAA
	(5473)																
Consensus																	
																ection	
	(5521)	5521		5	5530			554	0		55	550					5568
ClareAJ251507	(5214)	GCT	GGA	TTC	FATO	GAC.	ATG'	TTC	AAC	TTT	GAG	ACC'	TTT	GGC.	AAC	CAGC	ATG
huNaIII18 (AK)																	
	(5521)																
Consensus																	
																ection	
	(5569)	5569			,55				5590		_	560					5616
ClareAJ251507																	
huNaIII18 (AK)	(5105)	ATC	rgc:	rrg	rrc	CAA	ATT	ACA	ACC	TCT	GCT	GGC	TGG	GAT	GG	ATTG	CTA
JeongAF225987	(5569)	ATC	rgc:	rTG	rrc	CAA	АТТ	ACA	ACC	TCT	GCT	GGC	TGG	GAT	GG	ATTG	CTA
Consensus	(5569)	ATC	TGC	rTG:	rrc	CAA	ATT	ACA	ACC	TCT	GCT	GGC	TGG	GAT	GGZ	ATTG	CTA
															_	ection	
ClareAJ251507	(5617)	5617				563	30		5	640			5650	)			5664
ClareAJ251507	(5310)	GCA	CCT	TTA	CTT	TAA	AGT	GCA	CCA	CCC	GAC	TGT	GAC	CCT	GAG	CACA	TTA
huNaIII18 (AK)	(5153)	GCA	CCT	TTA	CTT	TAA	AGT	GCA	CCA	CCC	GAC	TGT	GAC	CCT	GAG	CACA	TTA
JeongAF225987	(5617)																
Consensus																	
																ection	
	(5665)	5665	.56	370			5680			569	90		.5	700			5712
ClareAJ251507	(5358)	CAC	CCT	GGC	AGC				GGA			GGG			TC	rgrī	
huNalli18 (AK)																	
JeongAF225987	(5665)																
Consensus																	
	( )																

									Section 120
(57 ClareAJ251507 (54	713)	5713	5720		5730	,5	740	5750	5760
ClareAJ251507 (54	406)	ATTTTC	CTTTTT	CGTCAG	TTACA	TCATCA	TATCCT	TCCTGG	TTGTGGTG
huNalli18 (AK) (52	249)	$\mathbf{A}\mathbf{T}\mathbf{T}\mathbf{T}\mathbf{T}\mathbf{C}$	TTTTT	<b>T</b> GTCAG	TTACA	TCATCA	TATCCT	TCCTGG	PTGTGGTG
JeongAF225987 (57	713)	TTTTA	CTTTTT	GTCAG	TTACA	TCATCA	TATCCT	TCCTGG	PTGTGGTG
Consensus (57	713)	TTTTA	TTTTTT	TGTCAG	TTACA	TCATCA	TATCCT	TCCTGG	TTGTGGTG
									Section 121
(57	761)	5761	577	00	<u>;</u> 5780		5790		5808
ClareAJ251507 (54	454)	AACAT	TACAT	CGCGGT	CATCC	TGGAGA	ACTTCA	GTGTTG	CTACTGAA
huNall118 (AK) (52	297)	AACATO	STACAT	CGCGGT	CATCC	TGGAGA	ACTTCA	GTGTTG	CTACTGAA
JeongAF225987 (57	761)	AACATO	STACAT	CGCGGT	CATCC	TGGAGA	ACTTCA	GTGTTG	CTACTGAA
Consensus (57	761)	AACAT	GTACAT	CGCGGT	CATCC	TGGAGA	ACTTCA	GTGTTG	CTACTGAA
									Section 122
. (58	809)	5809		5820	.58	330	5840		5856
ClareAJ251507 (55	502)	GAAAG	rgcaga	GCCCCT	GAGTG	AGGATG	ACTTTG	AGATGT'	TCTATGAG
huNaIII18 (AK) (53	345)	GAAAG	rgcaga	GCCCCT	GAGTG	AGGATG	ACTTTG	AGATGT	тстатсас
JeongAF225987 (58	809)	GAAAG'	rgcaga	GCCCCT	GAGTG	AGGATG	ACTTTG	AGATGT	TCTATGAG
Consensus (58	809)	GAAAG'	rgcaga	GCCCCT	GAGTG	AGGATG	ACTTTG	AGATGT	OZZOZZZOZ OZ OTATOT
									Section 123
(58	8571	5857		5870		5880	5	890	5904
ClareAJ251507 (55			AAAAS		TCCCG				
huNallI18 (AK) (53	393)	GTTTG	GAAAA	GTTTGE	TOCCG	ATGCGA ATGCGA	CCCAGT	TTATAG	$\mathbf{A}$ C $\mathbf{D}$ $\mathbf{D}$ $\mathbf{C}$ $\mathbf{D}$ $\mathbf{C}$ $\mathbf{D}$ $\mathbf{C}$
JeongAF225987 (58	857)	GTTTG	GAAAA	$G$ $\Psi$	TCCCG	ATCCCA ATCCCA	CCCAGT	TIAIAG.	AGTTCTCT AGTTCTCT
Consensus (58	857)	GTTTG	GAAAA	G $T$	TCCCG	A TROCA	CCCAGT	T T A T A G.	7 C
						2110007	CCCAGI		Section 124
(59	905)	5905	5910	,592	20	5930	١	5940	5952
ClareAJ251507 (55	598)	AAACT	CTCTGA	<u>, го</u>	AGCTG	CCCTGG	ATCCTC	CTCTTC	TCATAGCA
huNaIII18 (AK) (54	441)	AAACT	$^{\circ}$ T C T C I A	ጥጥጥጥርር	'AGCTG	CCCTGG	ATCCIC	CTCTTC	TCATAGCA
JeongAF225987 (59	905)	AAACT	$^{\circ}$	ጥጥጥጥርር	'AGCTG	CCCTGG	ATCCTC	CTCTTC	TCATAGCA
Consensus (59	905)	AAACT	CTCTGA	արարանշ	'AGCTG	CCCTGG		CTCTTC	TCATAGCA TCATAGCA
	/								Section 125
(50	9531	5953	5960		5970	5	980	5990	6000
ClareAJ251507 (56	646\	AAACC	777777	NCTCCT	CCMMA	mmccc r	A B C C A B C	<u>0666</u>	3000 30 4 0 80 0 8
huNaIII18 (AK) (54	48Q\	AAACC	CAACAA	AGTCCA		TIGCCP	A T G G W T C	TGCCCA	T GGT CAGT
JeongAF225987 (59	953\	AAACC	CAACAA	ACTCCA ACTCCA	CCLLY	mmcccr	MCCAMC	TGCCCA	TGGTCAGT TGGTCAGT
Consensus (59	9531	AAACC	CAACAA	ACTOCA		mmccc2	TGGATC	TGCCCA	TGGTCAGT
	000)	AAACC	CAACAA	MGICCA	GCIIA	TIGCCF	TIGGATC		- Section 126
161	004)	6001	604		0000		2000		
(b)	001)	6001	,601	00105	6020		6030		6048
ClareAJ251507 (50	094)	GGTGA	CCGGAT	CCACTO	FICTTG	ATATTI	'TATTTG	CCTTTA	CAAAGCGT
huNalli18 (AK) (5	004	GGTGA	CCGGAT	CCACTO	TCTTG	ATATTI	TATTTG	CCTTTA	CAAAGCGT
JeongAF225987 (6)	001)	GGTGA	CCGGAT	CCACTO	TCTTG	ATATTI	TATTTG	CCTTTA	CAAAGCGT
Consensus (6	UU1)	GGTGA	CCGGAT	CCACTO	FTCTTG	ATATT	TATTTG	CCTTTA	CAAAGCGI

									<del></del>						ion 127
(6 ClareAJ251507 (5	3049)	6049			,6060	)		6070	)		608	0			6096
ClareAJ251507 (5	5742)	GTTTT	G	GTG#	AGAG	TGG	AGA	GATG	GAT	GCC	CTT	CGA	ATAC.	AGAT	rggaa
huNalll18 (AK) (5	5585)	GTTTI	GQ(	GTG	AGAG	STGG	AGA	GATG	GAT	GCC	CTT	CGA	ATAC.	AGAT	rggaa
JeongAF225987 (6	3049)	GTTT	GT	GTG <i>I</i>	AGAG	TGG	AGA	GATG	GAT	GCC	СТТ	CGA	ATAC.	AGA:	rggaa
Consensus (6	3049)	GTTTT	rgg	GTG	AGAG	FTGG	AGA	GATG	GAT	GCC	CTT	CGA	ATAC.	AGA:	rggaa
															ion 128
	3097)					3110			120			6130			6144
ClareAJ251507 (5	5790)	GACA	GT	TTA	rggc	CATC	AAA	cccc	TCC	AAA	GTC	TCT'	PATG.	AGC	CTATT
huNalli18 (AK) (5	5633)	GACA	GT	TTA	rggc	CATC	AAA	cccc	TCC.	AAA	GTC	TCT	ratg.	AGC	TTATT
JeongAF225987 (6	6097)	GACA	GGT	TAT	rggc	CATC	AAA	cccc	CTCC	AAA	GTC	TCT	PATG.	AGC	CTATT
Consensus (6	6097)	GACA	GT	TTA	rggo	CATC	AAA	cccc	TCC	AAA	GTC	TCT	TATG.	AGC	CTATT
															ion 129
(6	6145)	6145	615	0		616	60		617	0		,61	80		6192
ClareAJ251507 (	5838)	ACAA	CCA	CTT	rga <i>i</i>	AACG	TAA	ACAF	AGAG	GAG	GTG	TCT	GCCG	CTA	TCATT
huNall118 (AK) (5	5681)	ACAA	CCA	CTT	rga <i>i</i>	AACG	TAA	ACA	AGAG	GAG	GTG	TCT	GCCG	CTA	ТСАТТ
															TCATT
Consensus (6	6145)	ACAA	CCA	CTT	rga <i>i</i>	AACG	TAA	ACA	AGAG	GAG	GTG	TCT	GCCG	CTA'	TCATT
														- Sect	ion 130
(€	6193)	6193		6200			6210			6220	)		6230		6240
ClareAJ251507 (	5886)	CAGC	GTA.	ATT	TCAC	GATG	ТТА	TCTT	ГТТА	AAG	CAA	AGG	TTAA	AAA	ATATA
huNall118 (AK) (	5729)	CAGC	GTA.	ATT	rcad	GATG	TTA	тстт	ГТТА	AAG	CAA	AGG	TTAA	AAA.	ATATA
JeongAF225987 (	6193)	CAGC	GTA.	ATT	TCA	GATG	TTA	TCT	TTA	AAG	CAA	AGG	AATT	AAA.	ATATA
Consensus (															
														- Sect	tion 131
(0	6241)	6241		,62	250		6	260		,e	270				6288
ClareAJ251507 (	5934)	TCAA	GTA	ACT.	ATA	ACAA	AGA	GGC	TTAA	AAA	GGG	AGG	ATTG	ACT	TACCT
huNallI18 (AK) (															
JeongAF225987 (	6241)	TCAA	GTA	ACT.	ATA	ACAA	AGA	GGC	ТТАА	AAA	GGG	AGG	ATTG	ACT	TACCT
Consensus (	6241)	TCAA	GTA	ACT.	ATA	ACAA	AGA	GGC	TTAA	AAA	AGGG	AGG	ATTG	ACT	TACCT
														- Sect	tion 132
(1	6289)	6289			630	0		631	0		632	20			6336
ClareAJ251507 (	5982)	ATAA	AAC	AAG	ACA'	TGAT	TAT	TGA	CAAA	CTA	TAAF	'GGG	AACT	CCA	CTCCA
huNall118 (AK) (	5825)	ATAA	AAC	AAG.	ACA'	TGAT	TAT	TGA	CAAA	CTA	raa <i>f</i>	GGG	AACT	CCA	CTCCA
															CTCCA
Consensus (	6289)	ATAA	AAC	AAG	ACA	TGAT	TAT	TGA	CAAA	CTA	raa <i>r</i>	'GGG	AACT	CCA	CTCCA
															tion 133
(	(6337)	6337			,	6350		9	6360			6370			6384
ClareAJ251507 (	(6030)	GAAA	AAA	CAG	ATG	GGAC	TTC	CTC	TACC	CAC	CICT	CCT	CCTI	CCT	ATGAT
huNallI18 (AK) (	(5873)	GAAA	AAA	CAG	ATG	GGAC	эттс	CTC	TACC	AC	CICI	CCT	CCTT	CCT	ATGAT
JeongAF225987 (	(6337)	GAAA	AAA	CAG	ATG	GGAC	TTC	CTC	TACC	CAC	ccci	CCT	CCTI	CCT	ATGAT
Consensus (															
`	•														

										<ul> <li>Section</li> </ul>	n 134
(638	5) <u>638</u>	5 ,63			100		6410		6420		6432
ClareAJ251507 (607	8) AG	TGTA	CAAA.	ACCAG	ACAAC	GAAA	AGTTT	GAGA	AGACA	AACC	AGAA
huNaIII18 (AK) (592	1) AG	TGTA	ACAAA.	ACCAG	ACAAC	GAAA	AGTTT	GAGA	AGACA	AACC	AGAA
JeongAF225987 (638	5) AG	TGTA	ACAAA.	ACCAG	ACAAC	GAAA	AGTTT	GAGA	AGACA	AACC	AGAA
Consensus (638	5) AG	TGTA	ACAAA.	ACCAG	ACAA	GAAA	AGTTT	GAGA	AGACA	AACC	AGAA
										_ Sectio	
(643	3) 643	3	6440		6450		6460	ı	6470	)	6480
ClareAJ251507 (612	6) AA	AGAA	AGCAA	AGGAA	AAGA	GTCA	GAGAA	AATCA	AAAAGI	AAAA	AGAA
huNall118 (AK) (596	9) AA	AGAA	AGCAA	AGGAA	AAGA	GTCA	GAGAA	AATC	AAAGI	CAAAA	AGAA
									AAAGT		
Consensus (643	3) AA	AGAA	AGCAA	AGGAA	AAGA	GTCA	GAGAA	AATC	AAAGT	CAAAA	AGAA
	<u>·</u>									Section	n 136
(648	1) 648	31	649		65	00	,6	510			6528
ClareAJ251507 (617	4) AC	AAAG	AATTA	TCTTT	GTGA	rcaai	TGTTI	ACAGO	CTATO	SAAGG'	TAAA
huNalli18 (AK) (601	7) AC	AAAG	ATTA	TCTTT	GTGA	rcaai	TGTTT	ACAG	CTATO	GAAGG'	ГААА
									CTATO		
Consensus (648											
								·		- Section	
(652	9) 652	29		6540		6550		6560			6576
ClareAJ251507 (622	2) GT	TATA			GACT		AGGAC	ance.	Mece	NA AVEN	
huNall118 (AK) (606	5) GT	TATA	GTGTC	AACTG	GACT	rcaac					
									ATGOO		o Kon
Consensus (652	9) GT	TATA	GTGTC	AACTG	GACT	rcaac	AGGAG	GTCC	ATGCC	AAACT	GACT
										_ Section	
(657	7) 657	77		6590		,66	00	66	10		6624
ClareAJ251507 (627	(O) (S)	TITA	ACKAA	TATER	ATAG	PERCY	Contract of the second	MANA	ATTACAT	enra a	THE R
huNallI18 (AK) (609	90)										
			Carlo Val	TAR THE	SAVIA CO	I SACE	THE COLUMN	1337 ha	NO NO A	A AND THE	
Consensus (657	7) GT	TTTA	ACAAA	TACTO	ATAG	TCAG	GCCTA	TACA	AGACA	GTGAA	GTGA
										- Section	
(663	25) 662	25 66	630	6	640		6650		6660		6672
ClareAJ251507 (631	18) <b>8</b>	UNITED IN		OMMO .	A TOMA			rerena.	TO COURT OF THE SECOND	A CHARLES	MACK ST
huNallI18 (AK) (609	30)		and the second								PARKET P
JeongAF225987 (662	25) 188	A CONTROL				TATE OF	l telefally	er estat v	CHAT		6.50
Consensus (662	25) CC	ጋጥጋጥ	<b>ТСТСР</b>	CTGC	АСПС	TCTC	AACCAC	CCTA	TCAAC	TTCA	CDDC
	-0, 00	-1010	10101		MCIC	10101	MOCH	JGGIA	LCAAC		on 140
299)	73) 66	73	6680		6690		670	n	671		6720
ClareAJ251507 (636 huNallI18 (AK) (609	コロノ コロノ (数数)							No Charles	HANGE S		
								Market Market			
									AGGAG		
Consensus (66)	(3) AC	. G. T. T. G	CTGTT	TTT.W.	TACC	AGCT	GACAC!	rgCTG	AGGAG.	AAACC	CAAT

	67211	6721		6720		2740		750		Section 14
ClareAJ251507 ( huNaIII18 (AK) ( JeongAF225987 ( Consensus (	6414)	Carro	a contract to	CA CHEAN	Mercial Company	3740	COMMON	750	VACOS SCHOOL	676
huNallI18 (AK) (	6090)								<b>MN D M</b>	
JeongAF225987 (	6721)	हाहार है।	ACCUA	GACTAT	AGGG	atiacoti	CRUCA	AAOUG	ACAMO	GTAACT
Consensus (	6721)	GGCT	ACCTA	GACTAT	AGGG	ATAGTI	GTGCA	AAGTG	AACATI	GTAACT
										Section 14
(	(6769)	6769		,6780		6790		6800		68
ClareAJ251507 (	6462)	CACC	AAACA	CCTT##	CTAC	AGTCCI	TGCAT	CCATT	TATTO	TTAACT
huNalli18 (AK) (	6090)	hunemann								
ClareAJ251507 ( huNallI18 (AK) ( JeongAF225987 (	6/69)	CACC	AAACA	CCTTT.	GTAC	agreci	TGCAT	CCATT	TATTO	TTAACT
Consensus (	0709)	CACC	AAACA	CCTTTA	AGTAC	AGTCCI	TGCAT	'CCATT(	CTATTI	TTAACT
	C047\	6017								Section 14
(Clare A 1251507	(6817) (6510)	001/		68	30	684	40	685	0	686
ClareAJ251507 ( huNallI18 (AK) ( JeongAF225987 (	(00 10)		1,112,113	S. C. A. LAN		AUDRICE	ANTI-CT	HE HAS	NC OF AIR	wite of the
JeongAF225987 (	6817)	le Real Print	Entering.		olarana.	CONTRACTOR OF	anyromy str	Dell'Article dell'		
Consensus (	6817)	CCAT	ATCTG	CCATAT	րդուրա	ACAAAA	ሳ ማ ማ ማ	$\Phi$	$\Gamma \subset C \setminus \Delta \cap \Gamma$	$\mathbf{T}$
	·							101110		Section 14
1	'A96E\	8888	6970		6000		6000	,	2000	00
ClareAJ251507 (	6558)	reco	CANTO	CATACI	THE TAX	TO VIEW	TOTAL	TUTCA	AA TO	m
ClareAJ251507 ( huNalli18 (AK) (	6090)									
JeongAFZZ5987 (	(6080	12 6 6	CONTRACT OF THE	CONTRACTOR	<b>医腹腔 基</b> 联	"是有事"第1日"管"	WHE CAND!	STEPHEN Y	CALL WITH	<b>Э</b> ТСТА А
Consensus (	(6865)	TCCC	CAATT	CATAGT	TATT	TCATAA	TGCTA	TGTCA	CTATTI	T
										Section 14
) ClareAJ251507 ( huNaIII18 (AK) (	(6913)	6913	692	20	693	0	6940	)	6950	696
CiareAJ251507 (	(6600)									
JeongAF225987 (	(6090)									
Consensus (	(6913) (6013)	TGAG	GTTTA	CGTTG	AAGAA.	ACAGT'A	TACAA	GAACC	CTGTCI	CTCAAA
										Section 14
	6961)	6961	_	6970		6980	6	000		
ClareAJ251507 ( huNalli18 (AK) (	(6600)							330	<del></del>	700
huNallI18 (AK) (	(6090)									
JeongAF225987 (	6961)	GATC	AGACA	AAGGTO	TTTT	GCCAGA	GAGAT	יתגגגגי	ኮጥጥጥርረ	CTCAAAA
Consensus (	(6961)									o i Chillin
		<del></del>								Section 14
(	(7009)	7009		,7020		,7030		,7040		70
ClareAJ251507 (	(6600)									
ClareAJ251507 ( huNaIII18 (AK) (	(6090)									
JeongAF225987 (	(7009)	CAGA	AAAAG	AATTGT	PAATG	GCTAC	AGTTTC	AGTTA	CTTCCA	ттттст
Consensus (	(7009)									

								Section 148
(70 ClareAJ251507 (66	057)	7057		,7070	,7080	,709	90	7104
huNaIII18 (AK) (60 JeongAF225987 (70 Consensus (70	090) 057)		GCTTTAA'		GTATTTTA	 GTCTGTTA	TGTTTG	TTTCTAT
(7)	105)	7105	7110	7120	.713	30	.7140	7152
(7 ClareAJ251507 (6 huNallI18 (AK) (6 JeongAF225987 (7 Consensus (7	090) 105)	CTGA	ACAGTTA		TAAAGTCT	CCTCTAAT	AATTTA	AGGATTA
(7	153)	7153						
(7 ClareAJ251507 (6 huNaIII18 (AK) (6 JeongAF225987 (7 Consensus (7	6600) 6090) 7153) 7153)	 TTTT		AGTATTCT				
(7 ClareAJ251507 (6 huNaIII18 (AK) (6 JeongAF225987 (7 Consensus (7	<sup>7</sup> 201)	TTTC	AGAGCTC	ATTTATAT	ATTTAGGT	CAAATGC	TTTCCA	AAAAGTAA
17	7240)	72/0		7260				
(7 ClareAJ251507 (6 huNalII18 (AK) (6 JeongAF225987 (7 Consensus (7	7249)	TCTA	АТАААТС	CATTCTAG	PATAAAAA	PATCTAAA	GTATTG	CTTTAGAA Section 153
(7	7297)	7297						
(7 ClareAJ251507 (6 huNaIII18 (AK) (6 JeongAF225987 (7 Consensus (7	6090) <mark>7297)</mark>	TAGT			TGCAGTA		CCATCT	TCTGCTCT
(7	7345)	7345	,7350	,7360				
(7 ClareAJ251507 (6 huNaIII18 (AK) (6 JeongAF225987 (7 Consensus (7	6090) 7345)	CAGO						

	/ <b>7</b> 000\	7000	7400	7440			Section 155
ClareAJ251507	(7393) (6600)	7393	,7400	,7410	7420	7430	7440
huNalil18 (AK)	(6090)						
JeongAF225987 Consensus	(7393)	TAGTI	PATTTTATCO	CTGTGGTGC	ATGTTTGGG	CAAATATAT	ATATAGCC
	(1000)		<del></del>				Section 156
,	(7441)	7441	7450	7460	74	70	7488
ClareAJ251507 huNaIII18 (AK)	(6600)						
huNall118 (AK)	(6090)						
JeongAF225987 Consensus	(7441) (7441)	TGATA	AAACAACTTC	TAAATTAATT	CAAATATGI	ACCACAGTG	TATGTGTC
	·				<del></del>		Section 157
ClareAJ251507 huNaIII18 (AK)	(7489)	7489	7500	75,	510	7520	7536
ClareAJ251507	(6600)						
huNaIII18 (AK)	(6090)						
JeongAF225987 Consensus	(7489)	TTTTC	GCAAGCTTC	CAACAGGGA	TGTATCCT	STATCATTCA:	TTAAACA1
Consensus	(7409)					· · · · · · · · · · · · · · · · · · ·	Section 158
	(7537)	7537					
ClareAJ251507	(6600)						
huNallI18 (AK)	(6090)						
JeongAF225987	/フにつづい	3 O m m a					
0	(1001)	AGTTT	PAAAGGCTA!	PCACTAATG	CATGTTAAI	CATTGCCTAT	GCTGCTCT
Consensus	(7537) (7537)	AGTTT					GCTGCTCT
Consensus	(7537)						GCTGCTCT  Section 159
Consensus	(7537)						GCTGCTCT  Section 159
Consensus ClareAJ251507 huNalil18 (AK)	(7537) (7585) (6600) (6090)	7585	,7590	,7600 	,7610 	,7620	GCTGCTC1 Section 159 7632
ClareAJ251507 huNalil18 (AK) JeongAF225987	(7537) (7585) (6600) (6090) (7585)	7585  ATTTS	,7590	,7600 	,7610 	,7620	GCTGCTC1 Section 159 7632
Consensus ClareAJ251507 huNalil18 (AK)	(7537) (7585) (6600) (6090) (7585)	7585  ATTTS	,7590 FACTCAATC	,7600 CATTCTTCA	,7610 	,7620 GGTTAAAGAA	GCTGCTCT Section 159 7632 TGTCACAT
ClareAJ251507 huNallI18 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585)	7585  ATTTT	,7590 	,7600  CATTCTTCA	,7610 	,7620 GGTTAAAGAA	GCTGCTCT Section 159 7632 TGTCACAT Section 160
ClareAJ251507 huNalII18 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585)	7585  ATTTT	,7590 	,7600  CATTCTTCA	,7610 	,7620 GGTTAAAGAA	GCTGCTCT Section 159 7632 TGTCACAT Section 160
ClareAJ251507 huNalII18 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585)	7585  ATTTT	,7590 	,7600  CATTCTTCA	,7610 	,7620 GGTTAAAGAA	GCTGCTCT Section 159 7632 TGTCACAT Section 160
ClareAJ251507 huNaIII18 (AK) JeongAF225987 Consensus ClareAJ251507 huNaIII18 (AK) JeongAF225987	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (6090) (7633)	7585 	,7590 FACTCAATCO ,7640	,7600 CATTCTTCA ,7650	,7610 CAAGTCTTC ,7660	,7620 GGTTAAAGAA'	Section 159 7632 TGTCACAT Section 160 7680
ClareAJ251507 huNall118 (AK) JeongAF225987 Consensus ClareAJ251507 huNall118 (AK)	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (6090) (7633)	7585 	,7590 FACTCAATCO ,7640 GTGATAGAA	,7600 CATTCTTCA 	,7610  CAAGTCTTC ,7660  CCTGCTCTC	,7620 GGTTAAAGAA ,7670 GTCCATTATG	Section 159 7632 TGTCACAT Section 160 7680
Consensus  ClareAJ251507 huNalil18 (AK) JeongAF225987 Consensus  ClareAJ251507 huNall118 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (6090) (7633) (7633)	7585 	,7590 FACTCAATCO ,7640 GTGATAGAA	,7600 CATTCTTCA ,7650 	,7610 CAAGTCTTC ,7660 	,7620 GGTTAAAGAA' ,7670 GTCCATTATG	Section 159 7632 TGTCACAT Section 160 7680 TCAAGCAC
ClareAJ251507 huNallI18 (AK) JeongAF225987 Consensus ClareAJ251507 huNallI18 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (6090) (7633) (7633)	7585 	,7590 FACTCAATCO ,7640 GTGATAGAA	,7600 CATTCTTCA ,7650 	,7610 CAAGTCTTC ,7660 	,7620 GGTTAAAGAA' ,7670 GTCCATTATG	Section 159 7632 TGTCACAT Section 160 7680 TCAAGCAC
ClareAJ251507 huNallI18 (AK) JeongAF225987 Consensus ClareAJ251507 huNallI18 (AK) JeongAF225987 Consensus	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (6090) (7633) (7633)	7585 	,7590 FACTCAATCO ,7640 GTGATAGAA	,7600 CATTCTTCA ,7650 	,7610 CAAGTCTTC ,7660 	,7620 GGTTAAAGAA' ,7670 GTCCATTATG	Section 159 7632 TGTCACAT Section 160 7680 TCAAGCAC
ClareAJ251507 huNaIII18 (AK) JeongAF225987 Consensus ClareAJ251507 huNaIII18 (AK) JeongAF225987	(7537) (7585) (6600) (6090) (7585) (7585) (7633) (6600) (7633) (7633) (7681) (6600) (6090) (7681)	7585 ATTTT  7633 ATTGC	,7590 FACTCAATCO ,7640 FTGATAGAA	_,7600 CATTCTTCA	,7610 CAAGTCTTC ,7660 CCTGCTCTC	,7620 GGTTAAAGAA ,7670 GTCCATTATG	GCTGCTCT Section 159 7632 TGTCACAT Section 160 7680 TCAAGCAC Section 161 7720

						Section	า 162
Clare A 10E4E07	(7729)	7729	,7740	,7750	,7760		7776
huNallI18 (AK)	(6000)						
JeongAF225987	(7729)	CAAC	ATGAGTATCATA	TGGTATCTC	гстссаттт		ים
Consensus	(7729)						
						Section	า 163
Clare A 1054507	(7777)	7777	7790	,780	0 ,78	10	7824
huNallI18 (AK)	(6090)						
JeongAF225987	(7777)	GGAT	ACTGCCTACTGA	CAAAACCTA	r	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	יים ב
Consensus	(7777)						
						Section	
Clara A 1054507	(7825)	7825	,7830 ,78	340	7850	,7860	787
huNall118 (AK)	(6090)						
JeongAF225987	(7825)	TGTC	FAAAACTTGTTT.	AAATATAAA	TAATGTAAA?		- — — ·
Consensus	(7825)						
						Section	n 165
Clara A 1254507	(7873)	7873	,7880 	,7890	,7900	,7910	792
huNall118 (AK)	(6090)						
JeongAF225987	(7873)	TATT	rgrcagcatttt	GTACATAAG.	TTATTAAAA	TTCAGGTTGAT	rga(
Consensus	(7873)						
						———— Section	n 166
Clare A 1251507	(7921)	7921	,7930 	7940	,7950		796
huNalli18 (AK)	(6090)						
JeongAF225987	(7921)	ATCA	CAATTTATTTA	CTTTATGCT	TTTGCTTTT	GATTTTTAATO	CAC
Consensus	(7921)						
						———— Section	n 167
Clara A 1254507	(7969)	7969	,7980	,7990	,8000		801
huNalli18 (AK)	(6090) (6090)						
JeongAF225987	(7969)	ATTC	CAAACTTTTGAA	TCCATAAGA	ТТТТТТТТТТТТТТТТТТ		מיחים:
Consensus	(7969)						
						Section	n 168
Olema A 1054507	(8017)	8017	,8030	,804	08, 0	50	808
huNall118 (ΔK)	(บบฮฮ) เกอกล)						
יייין טו וווואי ואוו	(5555)						
JeongAF225987	(8017)	AATA.	AAAGTTAGATAA	TGGGTTTTA	TGGATTTCT	<b>Γ</b> ΥGΥΥΑΥΑΑΥ	ላጥ ል
ClareAJ251507 huNall118 (AK)	(8017) (6600) (6090)	8017	8030	804	0 80		80

	(8065)	8065	8070	8080	8090	8100	Section 169
ClareAJ251507	(6600)					0100	
huNaIII18 (AK)	(6090)						
JeongAF225987 Consensus	(8065)	TTTC					
ClareAJ251507	(8113)	8113	8120	8130	8140	8150	8160
CialeAszs 1301	(OOOO)						
JeongAF225987	(8113)	ATCA	TTTTCTACCA	ACTATGGI	TGCCTCAAT	ATAACCTTT	ГАТТСАТА
Consensus	(8113)						
	(0161)	9161	9170	0400			Section 171
ClareAJ251507 huNaIII18 (AK)	(6600)		0170	,818,		90	8208
huNaIII18 (AK)	(6090)						
JeongAF225987	(8161)	GATG	TTTTTTTT	ATTCAACTI	TTGTAGTAT	TTACGTATG	CAGACTAG
Consensus	(8161)						0
	(8209)	8209	8220	) 8	230	8240	Section 172 8256
ClareAJ251507	(6600)						
huNall118 (AK)	(6090)						<b></b>
JeongAF225987 Consensus	(8209) (8209)	TCTT	ATTTTTTA	ATTCCTGCT	GCACTAAAG	CTATTACAA.	ATATAACA
	<u> </u>						Section 173
ClareAJ251507	(8257)	8257	8	3270	8280	,8290	8304
ClareAJ251507	(6600)						
JeongAF225987	(8257)	TGGA	 	 			
Consensus	(8257)	20011					
				<del></del>			Section 174
ClareAJ251507	(8305)	8305	,8310	8320	8330	8340	8352
huNalli18 (AK)	(6090)						
JeongAF225987	(8305)	TACC	TAACATGATA	rtttaaat.	TGTTTTTT	CACAAACCA	 AAAGTTTA
Consensus	(8305)						
	(0252)	9353	9360	9270	2000		Section 175
ClareAJ251507	(6600)	0353		83/0	8380	8390	8400
huNalli18 (AK)	(6090)						
JeongAF225987 Consensus	(8353)	ATGT	TAATTCTTT	TACAAAAC	TATTTACTO	STAGTGTATT	GAAGAACI
Consensus	(0353)						

							Sect	
(84 ClareAJ251507 (60	401)	8401	8410	,8420		_8430		8448
ClareAJ251507 (66 huNaIII18 (AK) (66	600)							
JeongAF225987 (84	401)	GCAT	GCAGGGAATT	GCTATTGC	 Фааааа	GAATGGTGA	A G C T A C G	 ጥሮልጥጥ
Consensus (84	401)	0 0111						
<del></del>								
(84 ClareAJ251507 (6	449)	8449	8460	8	470	8480		8496
huNaIII18 (AK) (6	0901							
JeongAF225987 (8	449)	ATTG.	AGCCAAAAGA	TTTAAATA	CATTTI	TTATTGCAT	TTCACT	TATTG
Consensus (8	449)					,		
(8- ClareAJ251507 (6-	497) 600)	8497	8:	010	8520	8530		8544
huNallI18 (AK) (6	(090)							
Inches A EQQEOQUE (0	4071	~~~	~~~~~~~					
Consensus (8	497)					~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		
(8 ClareAJ251507 (6	600)	8040	8550	,8560	8570	85	80	8592
huNallI18 (AK) (6 JeongAF225987 (8	090)							
JeongAF225987 (8	545)	TATA	ATAATTAATA	AAACCTGT	GCTTG	ATCTGACATT	<b>PTGTATA</b>	CATAA
Consensus (8	3545)						Coo	tion 100
(8	35931	8593						
(8 ClareAJ251507 (6	600)							
huNall118 (AK) (6	(090							
JeongAF225987 (8	3593)	AAGT	TTACATGAAT	TTTACAA	CAAACTA	AGTGCATGA:	TTCACCA	AGCAG
Consensus (8							Sec	tion 181
. (8	3641)	8641	,8650					
ClareAJ251507 (6	(0096				<del></del>			
huNallI18 (AK) (6								
JeongAF225987 (8 Consensus (8			ACAGAACAAA	GGCAAATT	PAAAAG	CAGCTTTGT	GAACTTT	TATGT
Consensus (d							Sec	tion 182
(8 ClareAJ251507 (6	3689)	8689						
ClareAJ251507 (6	3600)							
huNaIII18 (AK) (6	3090) 3690)	~~~~						
JeongAF225987 (8 Consensus (8			AAAGGATCAA	GTTCACA!	rGTTCC.	AACTTTCAG	GTTTGAT	AATAA
\$ 00.000.000 (C								

				F-1			- Section 183
() Clara & 1051507 (	(8737)	8737	<del></del>	8750	8760	8770	8784
huNallI18 (AK) (	(6090)						
huNallI18 (AK) ( JeongAF225987 ( Consensus (	(8737) (8737)	TAGT					
Claro A 1251507 (	(8785) (6600)	8785	8790	8800	8810	8820	8832
huNaIII18 (AK) (	(6090)						
JeongAF225987 ( Consensus (	(8785)	CTAT					
Olono A 1054507	(8833)	8833	,8840	885	0 8	860 887	0 888
						CATTCTTTGTT.	
Consensus (	(8833)						
	(0001)	8001	900		2000	0040	Section 186
ClareAJ251507 (	(6600)	0001			5900	8910	8928
huNallI18 (AK) (	(6090)					~~	
JeongAF225987 ( Consensus (	(8881) (8864)	CATT	ATATAAA	CTCCTATG	TATACATAA	GGTATTAATGA	TATAGTTAT
Consensus	(0001)						Section 187
(	(8929)	8929					
ClareAJ251507 (	(6600)	~				8960	
huNallI18 (AK) ( JeongAF225987 (	(6090) (8020)	ጥር አር	~~~~~~	~~~~~~			
Consensus (	(8929)	LGAG	MAITIAT.	ATTAACTT	TTTTTCAA	GAACCCTTGGA	TTTATGTGA
							— Section 188
Clare A 1054507.	(8977)	8977		8990	,9000	9010	9024
huNaiii18 (AK) (	(6090) (6090)	~					
JeongAF225987	(8977)	GGTC	AAAACCA	AACTCTTA	TTCTCAGTG	GAAAACTCCAG	TTGTAATGC
Consensus	(8977)						
	(0025)	0025				0000	
ClareAJ251507	( <i>9029)</i> (6600)	3023		3040	9050	9060	9072
huNalll18 (AK) (	(6090)						
JeongAF225987	(9025)	TATA	AAATTTT	GACAATTT	GGATCTAAA	TATGTATTTCA	TAATTCTCC
Consensus	(9025)						

					Sec	tion 190
(907	3) <u>9073</u>	,9080	,9090	9100	,9110	9120
ClareAJ251507 (660	0)					
huNaIII18 (AK) (609	0)					
JeongAF225987 (907	3) CATAA	TAAATTAT	ATAAGGTGGA.	AAAAAAAAA	AAAAAAAAA	AAAAA
Consensus (907						
					Sec	tion 191
(912	21) 9123			*		
ClareAJ251507 (660						
huNaIII18 (AK) (609	30)					
JeongAF225987 (912	1) AAA					
Consensus (912	21)					

							Section 1
	(1)		.10	20		30	40
ClareAJ251507protein	(1)	MAQALLV	PPGPES	FRLFTRES	LAAIE	EKRAAEEK	AKKPKKE
Translation of huNaIII18 (AK)	(1)	MAQALLV	PPGPES	FRLFTRES	LAAIE	EKRAAEEK	AKKPKKE
Translation of JeongAF225987	(1)	MAQALLV	PPGPES	FRLFTRES	SLAAIE	EKRAAEEK	AKKPKKE
Consensus	(1)	MAQALLV	PPGPES	FRLFTRES	LAAIE	EKRAAEEK	AKKPKKE
							- Section 2
	(41)	41	, <b>5</b> 0	60		.70	80
ClareAJ251507protein	(41)	QDNDDEN	KPKPNS	DLEAGKNI	PFIYO	GDIPPEMV	SEPLEDL
Translation of huNaIII18 (AK)	(41)	QDNDDEN	KPKPNS.	DLEAGKNI	PFIYO	GDIPPEMV	SEPLEDI
Translation of JeongAF225987	(41)	QDNDDEN	IKPKPNS	DLEAGKNI	PFIYO	GDIPPEMV	SEPLEDL
Consensus	(41)	QDNDDEN	KPKPNS	DLEAGKNI	PFIYO	DIPPEMV	SEPLEDL
							Section 3
	(81)	81	.90	,10	0	.110	120
ClareAJ251507protein			KTFIVM	NKGKAIFF			
Translation of huNaIII18 (AK)	(81)	DPYYINK	KTFIVM	NKGKAIFF	RESATS	SALYTLTP	T.NPVRKT
Translation of JeongAF225987	(81)	DPYYINK	KTFIVM	NKGKAIFF	RESATS	SALVII.TP	T.N.PVRKT
Consensus	(81)	DPYYINK	KTFIVM	NKGKAIFE	FSATS	SALVILUE SALVILUE	T.NDVPKT
					~	7775 2 2 11 2 1	- Section 4
	(121)	121	,130	,14	0	,150	- 00000011 160
ClareAJ251507protein				IMCTILTN	COLLENG	1100	OUT TOO
Translation of huNaIII18 (AK)	(121)	ATKTLVE	ISDI SML ISLESMI.	IMCTILTN		LLCNDDM	TKNVEXT
Translation of JeongAF225987	(121)	ATKTIVE	ISI FSMI.	IMCTILTI		MUGHENETI	TVNAFTT
Consensus	(121)	ATKTLVE	ISLESMI.	IMCTILTN	מאַעיבו <i>ע</i> דייענ מאַעיבו עדייענ		TUNNATION
	(			THEFT	CALLI	LUSMEEDW	Section 5
	(161)	161	,170	.18	n	100	•
ClareAJ251507protein				LARGFCLE		,190	200
Translation of huNaIII18 (AK)	(161)	エエジエエエ	TATTEST	LARGECLE LARGECLE	DEME	PDDPTMMT	DESVIVM
Translation of JeongAF225987	(161)	EUGIVUE	FSLIKI	LARGECLE	DEME	RDBMMMT	DESVIVM
Consensus	(161)	FTGITT	TATICES	LARGECLE LARGECLE	ים שמעני הדבד בחק	RDBMMMT	DESVIVM
	(,0,,		PDHIKI	THUGECUI	TOE LET		
	(201)	201	210	200	^		- Section 6
ClareAJ251507protein				22		230	240
Translation of huNall118 (AK)	(201)	AYVIEFV	SLGNVS	ALRTFRVI	RALKI	LISAIBGL	KTIVGAL
Translation of JeongAF225987	(201)	AYVORES	SLGNVS	ALRTFRVI	RALKI	risvipgL	KTIVGAL
Consensus	(201)	AIVTEEV	DLGNVS.	ALRTFRVI	RALKI	risvipgL	KTIVGAL
Consensus	(201)	AYVTEFV	SLGNVS.	ALRTFRVI	RALKI		
	(0.4.4)						<ul><li>Section 7</li></ul>
Olava & 105450**	(241)		250	26		270	280
ClareAJ251507protein		IQSVKKI	SDVMIL	TVFCLSVE	ALIGI	LQLFMGNL	RNKCLQW
Translation of huNaIII18 (AK)	(241)	IQSVKKI	SDVMIL	TVFCLSVE	ALIGI	LQLFMGNL	RNKCLQW
Translation of JeongAF225987	(241)	IQSVKKI	SDVMIL	TVFCLSVE	PALIGI	LQLFMGNL	RNKCLQW
Consensus	(241)	IQSVKKI	SDVMIL	TVFCLSVE	ALIGI	CQLFMGNL	RNKCLQW
A							

							Section 8
	281)	281	290		300	310	320
						FVNVTMST	
						TEMTVNVT	
						FVNVTMST	
Consensus (	(281)	PPSDSA:	FETNTT	SYFNGTI	MDSNGT	FVNVTMST	FNWKDYIG
							Section 9
,	(321)		330		,340	,350	360
ClareAJ251507protein (	(321)	DDSHFY	VLDGQK	DPLLCGI	NGSDAC	QCPEGYIC	VKAGRNPN
						GQCPEGYIC	
Translation of JeongAF225987 (	(321)	DDSHFY	VLDGQK:	DPLLCG	NGSDAC	GQCPEGYIC	VKAGRNPN
Consensus (	(321)	DDSHFY	VLDGQK:	DPLLCG	NGSDAC	GQCPEGYIC	VKAGRNPN
			~				<ul> <li>Section 10</li> </ul>
	(361)		370		380	390	400
ClareAJ251507protein (	(361)	YGYTSF	DTFSWA	FLSLFR:	LMTQDY	WENLYQLT	LRAAGKTY
						WENLYQLT	
						WENLYQLT	
Consensus (	(361)	YGYTSF	DTFSWA	FLSLFR:	LMTQD2	WENLYQLT	LRAAGKTY
							Section 11
	(401)		410		420	430	440
ClareAJ251507protein (	(401)	MIFFVL	VIFLGS	FYLVNL	ILAVVA	MAYEEQNQ	ATLEEAEQ
						AMAYEEQNQ	
						AMAYEEQNQ	
Consensus (	(401)	MIFFVL	VIFLGS	FYLVNL	ILAVVA	AMAYEEQNQ	ATLEEAEQ
							Section 12
	(441)		450		460	470	480
ClareAJ251507protein (	(441)	KEAEFQ	QMLEQL	KKQQEE.	AQAVA	AASAASRDF	SGIGGLGE
						ASAASRDF	
Translation of JeongAF225987 (	(441)	KEAEFQ	QMLEQL	KKQQEE.	AVAQA	AASAASRDF	SGGGGE
Consensus (	(441)	KEAEFQ	QMLEQL	KKQQEE.	AQAVA	AASAASRDF	SGIGGLGE
							Section 13
	(481)		490		_500	510	520
ClareAJ251507protein(	(481)	LLESSS	EASKLS	SK <b>S</b> AKE	WRNRRI	KKRRQREHL	EGNNKGER
						KKRRRREHL	
						KKRRQREHL	
Consensus (	(481)	LLESSS	EASKLS	SKSAKE	WRNRRI	KKRRQREHL	-
		•					Section 14
	(521)		530_		540	,550	560
ClareAJ251507protein	(521)	DSFPKS	ESEDSV	KRSSFL	FSMDGI	NRLTSDKKF	CSPHQSLL
						NRLTSDKKF	
						NRLTSDKKF	
Consensus	(521)	DSFPKS	ESEDSV	KRSSFL	FSMDGI	NRLTSDKKF	CSPHQSLL

				,		Section 15
	(561)	561	570	580	.590	600
				TSIFSFRGRAKI	OVGSENDF	ADDEHST
				TSIFSFRGRAKI		
Translation of JeongAF225987	(561)	SIRGSLFS	PRRNSK	TSIFSFRGRAKI	DVGSENDF	ADDEHST
				TSIFSFRGRAKI		
						Section 16
	(601)	601	610	620	630	640
ClareAJ251507protein			RDSLFVP	HRHGERRNS		
Translation of huNaIII18 (AK)				HRHGERRNS <b>NV</b>		MVPGLPA
Translation of JeongAF225987	(601)	FEDGESRI	RDSLFVP	HRHGERRNS <b>NV</b>	SQASMSSR	MVPGLPA
Consensus				HRHGERRNSNV		
						Section 17
	(641)	641	.650	.660	670	680
ClareAJ251507protein	(624)				N	GTTTETE
Translation of huNalII18 (AK)	(641)	NGKMHST	VDCNGVV	SLYGGPSALTS	PTGOLPPE	GTTTETE
Translation of JeongAF225987	(641)	NGKMHST	VDCNGVV	SLVGGPSALTS	PTGOLPPE	GTTTETE
Consensus	(641)	NGKMHST	VDCNGVV	SLVGGPSALTS	PTGOLPPE	GTTTETE
						Section 18
	(681)	681	690	.700	,710	720
ClareAJ251507protein				MLEDSSGRQRA		NTMEELE
Translation of huNaIII18 (AK)				MLEDSSGRQRA		
Translation of JeongAF225987				MLEDSSGRQRA		
Consensus				MLEDSSGRQRA		
						Section 19
	(721)	721	730	.740	,750	760
ClareAJ251507protein				NVFLIWDCCDA	WLKVKHLV	NLIVMDE
Translation of huNaIII18 (AK)				NVFLIWDCCDA		
Translation of JeongAF225987				NVFLIWDCCDA		
Consensus				NVFLIWDCCDA		
						Section 20
	(761)	761	.770	.780	.790	800
ClareAJ251507protein	(712)	FUDLATT		CLFMAMEHYPMI		VGNLVFT
Translation of huNall118 (AK)				CLFMAMEHYPMI		
Translation of JeongAF225987				LEMAMEHYPMI		
Consensus				CLEMAMEHYPMI		
						Section 21
	(801	801	.810	820	830	840
ClareAJ251507protein	•			MDPYYYFQEGWN		
Translation of huNaIII18 (AK)	•			MDPYYYFQEGW1		
Translation of JeongAF225987				MDPYYYFQEGW1		
Consensus				MDPYYYFQEGWI		
Consensus	(001	) GILIEDE	1 ^ 11 17 7 7 17 17	THE TATE OF OWN	,	~ ~ ~ ~ ~ (

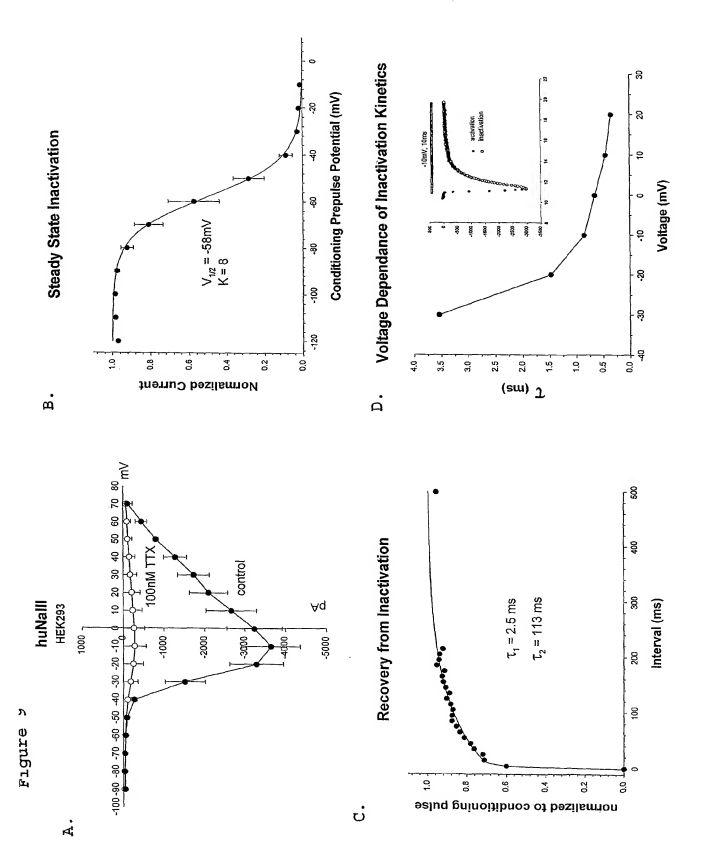
Section 24   Section 24   Section 25   Section 25   Section 25   Section 26   Section 26   Section 26   Section 26   Section 27   Section 27   Section 27   Section 27   Section 27   Section 27   Section 28   Sec					Se	ction 22
Translation of huNaill18 (AK)	(841)	841	850	860	870	880
Translation of JeongAF225987	ClareAJ251507protein (792)	LSNVEGLS	VLRSFRLLRV	FKLAKSWPTL	NMLIKI	GNSVG
Consensus   (841)   LSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVG		LSNVEGLS	VLRSFRLLRV:	FKLAKSWPTL	NMLIKI	GNSVG
Section 23   Section 23   Section 23   Section 23   Section 24   Section 24   Section of hundred in the section 24   Section 25   Section 25   Section 25   Section 26   Section 26   Section 26   Section 27   Section 27   Section 27   Section 27   Section 28   Section 29   Sec	Translation of JeongAF225987 (841)	LSNVEGLS	VLRSFRLLRV:	FKLAKSWPTL	NMLIKI	IGNSVG
Set   Set	Consensus (841)	LSNVEGLS'	VLRSFRLLRV:	FKLAKSWPTL	NMLIKI	IGNSVG
ClareAJ251507protein   Translation of huNallI18 (AK)   (81)   ALGNLTLVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT   (881)   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC   (921)   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC   (					Se	ection 23
Translation of huNalII18 (AK)   (881)   ALGNLTLVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT   (881)   ALGNLTLVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT   Consensus   (881)   ALGNLTLVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT   Section 24   Section 25   Section 26   Section 26   Section 26   Section 26   Section 25   Section 26   Section 27   Section 28   S		881	890	900	910	920
Consensus	ClareAJ251507protein (832)	ALGNLTLV	LAIIVFIFAV	VGMQLFGKSY	KECVCK	INDDCT
Consensus   (881)   ALGNLTLVLATIVFIFAVVGMQLFGKSYKECVCKINDDCT		ALGNLTLV	LAIIVFIFAV	VGMQLFGKSY	KECVCK:	INDDCT
Section 24   Section 25   Section 26   Section 27   Section 28   Sec	Translation of JeongAF225987 (881)	ALGNLTLV	LAIIVFIFAV	VGMQLFGKSY	KECVCK	INDDCT
Section 25   Section 25   Section 26   Section 27   Section 27   Section 28   Section 27   Section 28   Sec	Consensus (881)	ALGNLTLV	LAIIVFIFAV	VGMQLFGKSY	KECVCK	INDDCT
ClareAJ251507protein   Translation of huNallI18 (AK)   ClareAJ251507protein   ClareAJ2515					Se	ection 24
Translation of huNallI18 (AK) (921)	(921)	921	930	940	950	960
Consensus   1921   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC   1921   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC   1921   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC   1922   Section 25   1960   1970   1980   1990   1000	ClareAJ251507protein (872)	LPRWHMND	FFHSFLIVFR	VLCGEWIETM	WDCMEV	AGQTMC
Consensus   (921)   LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTMC	Translation of huNaIII18 (AK) (921)	LPRWHMND	FFHSFLIVFR	VLCGEWIETM	WDCMEV	AGQTMC
Section 25	Translation of JeongAF225987 (921)	LPRWHMND	FFHSFLIVFR	VLCGEWIETM	WDCMEV	AGQTMC
(961)   961   970   980   990   1000	Consensus (921	) LPRWHMND	FFHSFLIVFR	VLCGEWIETM	WDCMEV.	AGQTMC
ClareAJ251507protein Translation of huNallI18 (AK) Translation of JeongAF225987  (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Section 26  (1001) 1001					Se	ection 25
ClareAJ251507protein Translation of huNall118 (AK) Translation of huNall118 (AK) Translation of JeongAF225987 (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE Section 26 (1001) 1001	(961	961	970	.980	,990	1000
Translation of huNalII18 (AK) (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE  Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE  Consensus (961) LIVFMLVMVIGNLVVLNLFLALLLSSFSSDNLAATDDDNE  Section 26  (1001) 1001 1010 1020 1030 1040  ClareAJ251507protein (952) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Translation of huNalII18 (AK) (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Section 27  (1041) 1041 1050 1060 1070 1080  ClareAJ251507protein (992) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081			VIGNLVVLNL	FLALLLSSFS	SDNLAA	TDDDNE
Translation of JeongAF225987						
Section 26						
(1001) 1001	Consensus (961	) LIVFMLVM	VIGNLVVLNL	FLALLLSSFS	SDNLAA	TDDDNE
ClareAJ251507 protein (952) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Translation of huNall118 (AK) (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Translation of JeongAF225987 (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Section 27  (1041) 1041					S	ection 26
ClareAJ251507 protein (952) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Translation of huNall118 (AK) (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Translation of JeongAF225987 (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1041) 1041	(1001	) 1001	,1010	,1020	,1030	1040
Translation of huNall118 (AK) (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Translation of JeongAF225987 (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Section 27  (1041) 1041				KNKMRECFQK	AFFRKP	KVIEIH
Translation of JeongAF225987 (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Section 27  (1041) 1041						
Consensus (1001) MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH  Section 27  (1041) 1041 1050 1060 1070 1080  ClareAJ251507protein (992) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Translation of huNali18 (AK) (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081 1090 1100 1110 1120  ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF  Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF	Translation of JeongAF225987 (1001	) MNNLQIAV	GRMQKGIDYV	KNKMRECFQK	AFFRKP	KVIEIH
(1041) 1041 ,1050 ,1060 ,1070 1080 ClareAJ251507protein (992) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV Translation of huNall18 (AK) (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081 ,1090 ,1100 ,1110 1120  ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF						
ClareAJ251507protein (992) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV Translation of huNall18 (AK) (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081					S	ection 27
Translation of huNall18 (AK) (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081	(1041	) 1041	,1050	,1060	,1070	1080
Translation of huNali18 (AK) (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081	ClareAJ251507protein (992	) EGNKIDSC	MSNNTGIEIS	KELNYLRDGN	GTTSGV	GTGSSV
Translation of JeongAF225987 (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Consensus (1041) EGNKIDSCMSNNTGIEISKELNYLRDGNGTTSGVGTGSSV  Section 28  (1081) 1081 1090 1100 1110 1120  ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF  Translation of huNall18 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF  Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF						
ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of huNall18 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF	Translation of JeongAF225987 (1041	) EGNKIDSC	MSNNTGIEIS	KELNYLRDGN	GTTSGV	GTGSSV
ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of huNall18 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF						
ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of huNall118 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF					s	ection 28
ClareAJ251507protein (1032) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of huNall118 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF	(1081	) 1081	1090	,1100	,1110	1120
Translation of huNaill18 (AK) (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF						
Translation of JeongAF225987 (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF						
Consensus (1081) EKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEF	Consensus (1081	) EKYVIDEN	DYMSFINNPS	OVAIGVTVTLE	ESDFEN	LNTEEF

				S	ection 29
(1121)	1121	1130	,1140	,1150	1160
ClareAJ251507protein (1072)	SSESELE	ESKEKLNA	TSSSEGSTVDV	VLPREGEQ	AETEPE
Translation of huNall118 (AK) (1121)	SSESELE	ESKEKLNA	TSSEGSTVDV	VLPREGEQ	AETEPE
Translation of JeongAF225987 (1121)	SSESELE	ESKEKLNA	TSSSEGSTVDV	VLPREGEQ	AETEPE
Consensus (1121)	SSESELE	ESKEKLNA	TSSSEGSTVDV	VLPREGEQ	AETEPE
				S	ection 30
(1161)		,1170	,1180	,1190	1200
ClareAJ251507protein (1112)	EDLKPEA	CFTEGCIK	KFPFCQVSTEE	GKGKIWWN	LRKTCY
Translation of huNaIII18 (AK) (1161)	EDLKPEA	CFTEGCIK	KFPFCQVSTEE	GKGKIWWN	LRKTCY
			KFPFCQVSTEE		
Consensus (1161)	EDLKPEA	CFTEGCIK	KFPFCQVSTEE	GKGKIWWN	LRKTCY
	<del></del>			S	ection 31
(1201)		,1210	,1220	,1230	1240
ClareAJ251507protein (1152)	SIVEHNW	FETFIVFM	ILLSSGALAFE	DIYIEQRK	TIKTML
Translation of huNaIII18 (AK) (1201)	SIVEHNW	FETFIVFM	ILLSSGALAFE:	DIYIEQRK	TIKTML
Translation of JeongAF225987 (1201)	SIVEHNW	FETFIVFM	ILLSSGALAFE	DIYIEQRK	TIKTML
Consensus (1201)	SIVEHNW	FETFIVFM	ILLSSGALAFE	DIYIEQRK	TIKTML
					ection 32
(1241)		,1250	,1260	,1270	1280
ClareAJ251507protein (1192)	EYADKVF	TYIFILEM	ILLKWVAYGFQT	YFTNAWCW	LDFLIV
Translation of huNaIII18 (AK) (1241)	EYADKVF	TYIFILEM	ILLKWVAYGFQT	YFTNAWCW	LDFLIV
Translation of JeongAF225987 (1241)	EYADKVF	TYIFILEM	ILLKWVAYGFQT	YFTNAWCW	LDFLIV
Consensus (1241)	EYADKVF	TYIFILEM	ILLKWVAYGFQT	YFTNAWCW	LDFLIV
					ection 33
(1281)		1290	,1300	,1310	1320
ClareAJ251507protein (1232)	DASPASP	VANALGYS	SELGAIKSLRTL	RALRPLRA	LSRFEG
Translation of huNaIII18 (AK) (1281)	DASTAST	VANALGYS	SELGAIKSLRTL	RALRPLRA	LSRFEG
Translation of JeongAF225987 (1281)	DASTAST	VANALGYS	ELGAIKSLRTL	RALRPLRA	LSRFEG
Consensus (1281)	DASTAST	VANALGYS	SELGAIKSLRTL		
				=	ection 34
(1321)		1330	,1340	,1350	1360
ClareAJ251507protein (1272)	MRVVVNA	LVGAIPSI	MNVLLVCLIFW	LIFSIMGV	NLFAGK
Translation of huNaIII18 (AK) (1321)	MRVVVNA	LVGAIPSI	MNATTACTIEM	LIFSIMGV	NLFAGK
			MNVLLVCLIFW		
Consensus (1321)	MRVVVNA	LVGAIPSI	MNVLLVCLIFW		
					ection 35
(1361)		,1370	,1380	,1390	1400
ClareAJ251507protein (1312)	FYHCVNM	TTGNMFD	SDVNNLSDCQA	LGKQARWK	NVKVNF
Translation of huNaIII18 (AK) (1361)	FYHCVNM	TTGNMFD	SDVNNLSDCQA	LGKQARWK	NVKVNF
			SDVNNLSDCQA		
Consensus (1361)	) FYHCVNM	TTGMMFDI	SDVNNLSDCQA	LGKQARWK	NVKVNF

					Section 36
(1401)	1401	1410	,1420	,1430	1440
ClareAJ251507protein (1352)				YAAVDSRDVK	LQPVYEE
Translation of huNall18 (AK) (1401)					
Translation of JeongAF225987 (1401)	DNVGAG	SYLALLQV	ATFKGWMDIM	YAAVDSRDVE	CLQPVYEE
Consensus (1401)					
					Section 37
(1441)	1441	,1450	,1460	,1470	1480
ClareAJ251507protein (1392)	NLYMYI	YFVIFII	FGSFFTLNLF	IGVIIDNFN	QKKKFGG
Translation of huNaIII18 (AK) (1441)	NLYMYI	YFVIFII	FGSFFTLNLF	IGVIIDNFN	QQKKKFGG
Translation of JeongAF225987 (1441)	NLYMYI	YFVIFII	FGSFFTLNLF	IGVIIDNFN	QQKKKFGG
Consensus (1441)	NLYMYI	LYFVIFII	FGSFFTLNLF	IGVIIDNFN(	QQKKKFGG
			<del></del>		Section 38
(1481)	1481	,1490	,1500	,1510	1520
ClareAJ251507protein (1432)	QDIFM	PEEOKKYY	NAMKKLGSKK	PQKPIPRPA	NKFQGMVF
Translation of huNall18 (AK) (1481)	QDIFM'	PEEQKKYY	NAMKKLGSKK	PQKPIPRPA	NKFQGMVF
Translation of JeongAF225987 (1481)	QDIFM	PEEQKKYY	NAMKKLGSKK	PQKPIPRPA	NKFQGMVF
Consensus (1481)	QDIFM'	TEEQKKYY	NAMKKLGSKK	PQKPIPRPA	NKFQGMVF
					_ Section 39
(1521)	1521	,1530	,1540	1550	1560
ClareAJ251507protein (1472)	DFVTR	QVFDISIM	ILICLNMVT	MVETDDQGK	YMTLVLSR
Translation of huNaIII18 (AK) (1521)	DFVTR	QVFDISIM	ILICLNMVTN	MVETDDQGK	YMTLVLSR
Translation of JeongAF225987 (1521)	DFVTR	QVFDISIM	ILLICLNMVT	MVETDDQGK	YMTLVLSR
Consensus (1521)	DFVTR	QVFDISIM	ILLICLNMVTN	MVETDDQGK	YMTLVLSR
					_ Section 40
(1561)	1561	1570	,1580	,1590	1600
ClareAJ251507protein (1512)	INLVF	IVLFTGEF	VLKLVSLRHY	YFTIGWNIF	DFVVVILS
Translation of huNaIII18 (AK) (1561)	INLVF	IVLFTGER	VLELVSLRHY	YYFTIGWNIF	DFVVVILS
Translation of JeongAF225987 (1561)	INLVF	IVLFTGE	VL <b>K</b> LVSLRH	YFTIGWNIF	DFVVVILS
Consensus (1561	INLVF	IVLFTGE	VLKLVSLRH	CYFTIGWNIF	DFVVVILS
					Section 41
(1601	1601	.1610	,1620	,1630	
ClareAJ251507protein (1552	IVGMF	LAEMIEKY	FVSPTLFRV:	IRLARIGRIL	RLIKGAKG
Translation of huNaIII18 (AK) (1601	, VIVGMF	LAEMIEKY	FVSPTLFRY:	IRLARIGRIL	RLIKGAKG
Translation of JeongAF225987 (1601	)   IVGMF	LAEMIEK	SVSPTLFRV:	IRLARIGRIL	RLIKGAKG
Consensus (1601	) IVGMF	LAEMIEK	YFVSPTLFRV:	IRLARIGRIL	RLIKGAKG
					_ Section 42
(1641	) 1641	1650	. ,1660	. 1670	1680
ClareAJ251507protein (1592	IRTLL	FALMMSL		FLVMFIYAIF	GMSNFAYV
Translation of huNaIII18 (AK) (1641					
			PALFNIGLLL		
Consensus (1641					
•	-				

				S	ection 43
(1681)	1681	,1690	1700	1710	1720
ClareAJ251507protein (1632)		MFNFETFGN	SMICLFQITT	SAGWDGL	LAPILN
Translation of huNall118 (AK) (1681)					
Translation of JeongAF225987 (1681)	KKEAGIDDI	MFNFETFGN	SMICLFOITT	SAGWDGI	LAPILN
Consensus (1681)					
					Section 44
(1721)	1721	1730	,1740	,1750	1760
ClareAJ251507protein (1672)	SAPPDCDP	DTIHPGSSV	KGDCGNPSVO	GIFFFVSY	ZIIISFL
Translation of huNall118 (AK) (1721)					
			KGDRGDPSV		
Consensus (1721)					
					Section 45
(1761)	1761	,1770	,1780	,1790	1800
ClareAJ251507protein (1712)	VVVNMYIA	VILENFSVA		EDDFEME	YEVWEKF
Translation of huNall118 (AK) (1761)					
			TEESAEPLS		
Consensus (1761)					
					Section 46
(1801)	1801	1810	,1820	,1830	1840
ClareAJ251507protein (1752)	DPDATOFT	EFSKLSDE	AAI,DPPI,I,T		LIAMDLP
Translation of huNall18 (AK) (1801)					
			AAALDPPLLI		
Consensus (1801)					
					Section 47
(1841)	1841	1850	,1860	,1870	1880
ClareAJ251507protein (1792)					
Translation of huNaIII18 (AK) (1841)					
			TKRVLCESGE		
Consensus (1841)					
					Section 48
(1881)	1881	,1890	,1900	.1910	1920
ClareAJ251507protein (1832)	SNPSKVSY				
Translation of huNall118 (AK) (1881)	SNISKVSI	EPITTTER	RKOEEVSAAT	TORNERC	VIIKORI
Translation of JeongAF225987 (1881)	SNICKVSI	EPTTTTLK	RKQEEVSAAI	TORNERC	VI.I.KORI.
Consensus (1881)					
					Section 49
(1034)	1921	,1930	,1940	.1950	1960
ClareAJ251507protein (1872	WNTCONV	INENTEGRE			
Translation of huNaIII18 (AK) (1921	NNT CCNVN NNT CCNVN	INDATEGET.	DI'DIKUDW <sub>I</sub> i PPETYÄDMTT	DKI'MGMG	LDEKLDG TIEKIDG
Translation of JeongAF225987 (1921	ANT CONTY Y WIT CONTY	INDAINGRI INDAINGRI	DIOTECTIONS	DKINGNG	TIEKIDG
Consensus (1921	Y WILGGMAN	TADATAGNI	DI.DIKUDMII	DKINGME	TIBRIDG
Consensus (1921	V WINT DOIN I	NUDAINGKI	PHETVANITT	CHEMITY	11111110

				Se	ction 50
(1961)	1961	1970	1980	,1990	2000
ClareA.1251507protein (1912)	SSSTT	PPSYDSVTK	PDKEKFEKDK	PEKESKGKEV	JRENQK
Translation of huNall118 (AK) (1961)	SSSTT	PPSYDSVTK	PDKEKFEKDK!	PEKESKGKE	JRENQK
Translation of JeongAF225987 (1961)	SSSTTI	PPSYDSVTK	PDKEKFEKDK	PEKESKGKE	VRENQK
Consensus (1961)	SSSTTS	SPPSYDSVTK	PDKEKFEKDK	PEKESKGKE	VRENQK



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- (71) Applicant (for all designated States except US): EURO-CELTIQUE S.A. [LU/LU]; 122, Boulevard de la Petrusse, L-2330 Luxembourg (LU).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): KAMMESHEIDT, Anja [DE/US]; 31558 Eagle Rock Way, Laguna Beach, CA 92651 (US). HODGES, Dianne [US/US]; 14351 Pinewood Road, Tustin, CA 92780 (US).
- Agents: ROBINSON, Joseph, R. et al.; Darby & Darby P.C., P.O. Box 5257, New York, NY 10150-5257 (US).

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(54) Title: SPLICE VARIANT OF HUMAN SODIUM III CHANNEL (HNAII118)

(57) Abstract: Described herein is a splice variant of the human NaIII channel α subunit, designated hNaIII18. Also described are nucleotide and amino acid sequence for hNaIII18, oligonucleotide primers and probes for hNaIII18, hNaIII18 regulatory sequences, hNaIII18-specific antibodies, methods of detecting hNaIII18 proteins or nucleic acids, and methods of screening for modulators of hNaIII18 expression or activity.

## INTERNATIONAL SEARCH REPORT

International application No.
PCT/US03/38796

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IPC	(7)	: C12 Q 1/68		- 04.01	ļ
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INTERNATIONAL SEARCH REPORT	PCT/US03/38796
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search terms: nucleotide, polypeptide, sodium channel, human.	
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